

Paramagnetic Chemical Probes for Studying Biological Macromolecules

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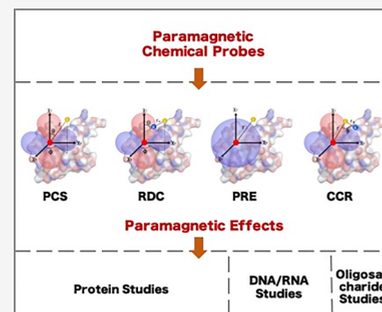
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ABSTRACT: Paramagnetic chemical probes have been used in electron paramagnetic resonance (EPR) and nuclear magnetic resonance (NMR) spectroscopy for more than four decades. Recent years witnessed a great increase in the variety of probes for the study of biological macromolecules (proteins, nucleic acids, and oligosaccharides). This Review aims to provide a comprehensive overview of the existing paramagnetic chemical probes, including chemical synthetic approaches, functional properties, and selected applications. Recent developments have seen, in particular, a rapid expansion of the range of lanthanoid probes with anisotropic magnetic susceptibilities for the generation of structural restraints based on residual dipolar couplings and pseudocontact shifts in solution and solid state NMR spectroscopy, mostly for protein studies. Also many new isotropic paramagnetic probes, suitable for NMR measurements of paramagnetic relaxation enhancements, as well as EPR spectroscopic studies (in particular double resonance techniques) have been developed and employed to investigate biological macromolecules. Notwithstanding the large number of reported probes, only few have found broad application and further development of probes for dedicated applications is foreseen.



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1. PARAMAGNETIC NMR FOR BIOMOLECULAR STUDIES

1.1. General Introduction

Nuclear magnetic resonance (NMR) spectroscopy is the most widely used spectroscopic technique to obtain structural information with atomic resolution. The NMR observable most frequently used for structure determination is the nuclear Overhauser effect (NOE), which measures dipolar interactions between protons separated by up to about 5 Å and, thus, provides internuclear distance restraints.^{1,2} Owing to the much larger magnetic moment of unpaired electrons compared with nuclei (about 658 times greater for a lone electron than for a proton), dipolar interactions involving unpaired electrons, such as in metalloproteins containing paramagnetic metal ions, can be detected over a much greater distance range. As the paramagnetic effects of unpaired electrons on the NMR spectrum tend to be very large and are readily detected, potentially over distances up to 100 Å from the paramagnetic center, they present valuable long-range structural information. In fact, 50 years ago and prior to the invention of two-dimensional NMR spectroscopy, paramagnetic effects were thought to be the key to 3D structure determinations of proteins and other biomolecules in solution.^{3,4} The paramagnetic effects are in many instances

anisotropic, contributing not only distance but also orientational restraints.

Apart from uses in NMR spectroscopy, paramagnetic labels also feature most prominently in electron paramagnetic resonance (EPR) spectroscopy, where they have gained in importance by their capacity to accurately measure distances between paramagnetic centers on the nanometer scale. This Review will focus on applications in NMR spectroscopy, while also acknowledging the performance of tags for distance measurements by EPR spectroscopy, specifically in double-electron–electron resonance (DEER) experiments. Furthermore, europium(III) and terbium(III) are paramagnetic metal ions which, in conjunction with an aromatic antenna group, can yield intense luminescence suitable for distance measurements by Förster resonance energy transfer (FRET).^{5,6} The same probes can thus be suitable for NMR and FRET applications, but luminescence applications are not discussed in this Review which focuses on paramagnetism and NMR spectroscopy.

The field of paramagnetic NMR has been reviewed before. In particular, the theory of paramagnetic NMR and the resulting spectral effects have been described in many reviews and books, covering magnetic susceptibility,^{7–11} relaxation,¹² and effects in solids¹³ in great detail. This Review presents an integrated formalism of the various effects and includes the most recent effects reported.

Initially, paramagnetic NMR spectroscopy of biological macromolecules took advantage of paramagnetic metal ions present in proteins,^{14,15} and this approach is still finding application for the study of metalloproteins and also in research into metal trafficking.¹⁶ Also metal substitution has been used extensively.¹⁷ With the arrival of paramagnetic tags, the application of the tool box of paramagnetic NMR spectroscopy became available for biomolecules that lack a natural paramagnetic center, expanding its potential greatly. Several reviews give a general overview of the possibilities of employing pseudocontact shifts (PCSs), paramagnetic relaxation enhancements (PREs), and residual dipolar couplings (RDCs) (terms to be explained below) or focus on the range of different tags available and their applications.^{18–26}

Specific applications have been the topic of dedicated reviews. One example is the use of paramagnetism for partial alignment to generate RDCs for the study of structure and dynamics.²⁷ PCSs provide useful data for the structure determination of proteins and protein–protein or protein–ligand complexes, as sole restraints, or in combination with other data, such as X-ray diffraction and small-angle X-ray scattering.^{10,26,28–30} PREs are useful in particular because of their ability to report on minor, “invisible” or “dark” states present in a sample.^{31–34} PREs generated by soluble probes, that is, paramagnetic molecules not covalently linked to the molecule of interest, are referred to as solvent PREs and can provide information on molecular surfaces and changes in surface exposure upon complex formation.^{35,36} A recent review describes the use of paramagnetic NMR in drug discovery.³⁷ Tags have been mostly designed for and applied to proteins, but applications to other biomolecules have started to appear, such as oligosaccharides,^{38,39} and a significant body of work exists describing oligonucleotides with paramagnetic centers for EPR measurements.^{40–47}

The main aim of this Review is to describe the state-of-the-art of paramagnetic tags for biological macromolecules. We wish to illustrate the diversity of tags and discuss their pros and cons for various applications. Contrary to most other reviews, particular attention is paid to the chemical synthesis of the tags because the

chemistry of the more advanced cyclen-based lanthanoid tags is not trivial and in some cases a limiting factor, due to demanding synthetic routes or limited chemical stability under physiological conditions and in the presence of proteins. Paramagnetic compounds have many more applications, e.g., as contrast agents in magnetic resonance imaging (MRI) and polarizing agents for dynamic nuclear polarization (DNP). These topics are considered outside the scope of this Review, and the reader is referred to excellent, recently published reviews.^{48,49} Paramagnetic tags attached to proteins exclusively for the purpose of studies by EPR spectroscopy have been comprehensively discussed previously.⁵⁰ Similarly, our discussion omits EPR spin labels attached via a long and flexible tether, such as resulting from tagging noncanonical amino acids.⁵¹ In contrast, our account attempts to give a comprehensive overview over paramagnetic tags designed specifically for structure studies of deoxyribonucleic acid (DNA) and ribonucleic acid (RNA) oligomers by NMR and EPR spectroscopy. Finally, various paramagnetic compounds have been devised as sample additives to enhance longitudinal relaxation rates in solid-state NMR spectroscopy.^{52,53} As far as these compounds are not chemically attached to biological macromolecules, the reader is again referred to a recent review for a more complete compilation.⁵⁴

As paramagnetic probes elicit a multitude of effects depending on their chemical, physical, and dynamic properties, an increasing number of probes have been synthesized with the aim of facilitating their installation in biological macromolecules and maximizing the information content that can be gathered from the observation of the paramagnetic effects. About half of the published paramagnetic probes were described for the first time in the past five years. To assess their utility for different purposes, it is necessary to understand the origin and manifestation of the paramagnetic effects. Many of the most stringent tag requirements stem from applications in paramagnetic NMR spectroscopy.

The terms spin label, paramagnetic tag and paramagnetic probe are frequently found in the literature and also used throughout this Review. They all refer to paramagnetic compounds that are introduced and used to generate paramagnetic effects to study a system of interest. In general, these terms can be considered synonyms. The term “spin label” is probably the oldest and has been widely used in the field of EPR for many years. It refers to labeling a system with an unpaired electron spin. “Paramagnetic tag” emphasizes the quality of a small label to give information about the bearer, that is, provide site-specific information about the tagged molecule. The term “paramagnetic probe” relates more generally to the function, that is, probing the molecule under investigation and includes also soluble compounds that are not bound to specific sites.

1.2. Qualitative Description of Paramagnetic Effects in NMR Spectroscopy

Paramagnetic effects detected in NMR spectra depend on the dipolar fields generated by unpaired electrons. If the electron spins relax slowly compared with the rotational correlation time of the electron–nucleus vector, the dipolar field of the electron at the site of the nuclear spin averages to zero, provided that the molecule reorients isotropically in solution. In this case, the chemical shift of the nuclear spin does not change, but the time fluctuation of the dipolar field can greatly enhance the nuclear relaxation. Electron spins that relax rapidly compared with the rotational correlation time of the electron–nucleus vector lead to a Curie spin, which is the time-averaged net magnetic

moment of the electron spin aligned with the external magnetic field. The Curie spin constitutes a molecular magnetic susceptibility. In this situation, it is useful to describe the magnetic susceptibility associated with the paramagnetic center, χ , by a tensor that defines its magnitude as a function of the molecular orientation in the external magnetic field. In general, the χ tensor of a paramagnetic center with rapidly relaxing electrons is anisotropic and its anisotropic component is commonly referred to as $\Delta\chi$ tensor.

Chemical shift changes observed in NMR spectra due to the presence of a paramagnetic center are referred to as hyperfine shifts. For paramagnetic centers with a Curie spin, hyperfine shifts comprise two parts, the contact shift and PCS. Contact shifts are a consequence of the delocalization of unpaired electron spin density across chemical bonds and thus are only observed for nuclear spins fairly close to the paramagnetic center. In contrast to the contact shift, the PCS is a through-space interaction, which arises from the time-averaged dipolar interaction between the unpaired electron spins and the nuclear spin. PCSs are observable over much greater distances and, as they are independent of bond angles, can be described by fewer parameters, which pertain to the $\Delta\chi$ tensor and the location of the nuclear spin relative to the $\Delta\chi$ tensor. In this way, PCSs deliver long-range distance and orientation information, which can readily be interpreted.

Anisotropic magnetic susceptibilities also cause weak alignment of the paramagnetic molecule in the external magnetic field, leading to the observation of RDCs between nuclear spins. In as far as the molecule tumbles in solution as a single rigid entity, the alignment affects the entire molecule irrespective of the location of the paramagnetic center and, therefore, RDCs do not depend on the distance of the nuclear spins from the paramagnetic center. Conveniently, the size and orientation of the alignment tensor are proportional to the size and orientation of the $\Delta\chi$ tensor.

A paramagnetic center with Curie spin generates a magnetic dipolar field at the site of a nuclear spin. The mathematical description of the effect of this dipolar field on the magnetic shielding of the nucleus is closely similar to that of chemical shift anisotropy (CSA). To highlight this similarity, the term dipolar shielding anisotropy (DSA) has been coined for this paramagnetic effect. The similarity extends to cross-correlated relaxation (CCR) effects between DSA and dipole–dipole (DD) relaxation, which is manifested in differential relaxation rates of multiplet components, just like the CSA/DD cross-correlation effects that form the basis of transverse relaxation optimized spectroscopy (TROSY).^{55,56} DSA/DD cross-correlated relaxation can be used to evaluate the angles between an internuclear bond and the principal axes of the DSA tensor. As the magnitude of the DSA tensor depends on the distance of the nuclear spin from the paramagnetic center, conversion of CCR effects into structural restraints is more involved than for RDCs. Nonetheless, these CCR effects are readily observable and can provide useful distinctions between different pairs of coupled nuclear spins that otherwise display similar paramagnetic effects.⁵⁷

A different type of CCR effect also occurs between CSA and DSA relaxation. This CCR effect can significantly affect the nuclear net relaxation rates, either enhancing or reducing the relaxation of nuclear magnetization in the paramagnetic state compared with the corresponding diamagnetic state.⁵⁸

In general, paramagnetic centers enhance the relaxation rates of nuclear spins, which is referred to as PRE. A PRE is the direct

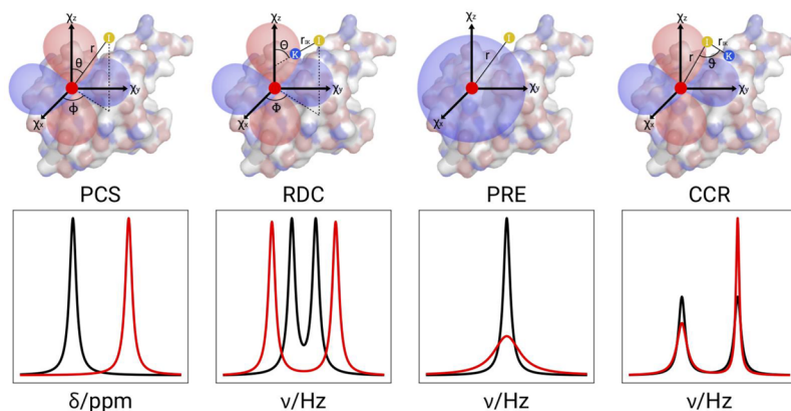


Figure 1. Schematic representation of the paramagnetic effects of PCS, RDC, PRE, and CCR, respectively, where CCR refers to cross-correlation between DSA and dipole–dipole relaxation. The top panel depicts the geometric dependencies relative to the frame of a χ tensor. The bottom panel illustrates the effects in the NMR spectra of paramagnetic (red) versus diamagnetic (black) samples.

consequence of dipole–dipole interactions between the paramagnetic center and a nuclear spin. It decreases with the sixth power of the distance between them. The large magnitude of the PRE at short distances and its steep distance dependence render the PRE a sensitive tool to detect minor states that position the nucleus close to the unpaired electrons, as may occur in dynamic proteins carrying a paramagnetic tag. Paramagnetic centers with slow electronic relaxation ($\sim 10^9$ s $^{-1}$), such as presented by nitroxide radicals, Mn(II), and Gd(III) ions, generate PREs by dipole–dipole relaxation, referred to as Solomon–Bloembergen–Morgan (SBM) or Solomon relaxation.⁵⁹ In contrast, SBM relaxation is much less efficient for unpaired electrons with fast electronic relaxation rates ($> 10^{11}$ s $^{-1}$) and Curie spin relaxation becomes more prominent,^{60,61} especially for long rotational correlation times of the molecule and at increased magnetic field strength. In general, the SBM mechanism is the sole contribution to PREs in isotropic paramagnetic species and the Curie spin mechanism adds an additional component in systems characterized by paramagnetic anisotropy, which can nonetheless become the dominant contribution to PREs.

Many transition metal ions contain unpaired electrons, leading to paramagnetism. The paramagnetism of 3d block and 4f block ions has been studied most thoroughly.^{29,62} Mn(II) ions have a long electronic relaxation time, causing no PCSs but strong PREs. High-spin Co(II) features a Curie spin and can generate large PCSs with weak PREs, compared with other 3d block ions. In aqueous solution, iron is stable in the oxidation states +2 and +3. Depending on its ligands, Fe(II) can be paramagnetic or diamagnetic. Similarly, the paramagnetism of Fe(III) can be either large (high spin) or small (low spin), depending on the coordination environment. The 4f block elements are referred to as lanthanoids,⁶³ with the general symbol Ln. Unlike 3d block ions, lanthanoids are seldom involved in biological processes, with the first examples of lanthanoenzymes discovered only in 2011 in methanotrophic bacteria.⁶⁴ Lutetium(III) and lanthanum(III) have no unpaired electrons, while all other Ln(III) ions of the series are paramagnetic. Gd(III) ions are unique for featuring a strong isotropic magnetic susceptibility, which is generated by seven unpaired electrons. Gd(III) ions yield exceptionally strong PREs without generating PCSs. Other Ln(III) ions have Curie spins of different magnitude, causing both PCS and PRE effects. Stable organic radicals such as nitroxides or the trityl radical can also be

used to create a paramagnetic center. They do not cause PCSs, but yield sizable PREs.

Quantification of the paramagnetic effects is commonly achieved by comparison with a diamagnetic species. Therefore, the appropriate choice of a diamagnetic reference as similar to the paramagnetic sample as possible is very important. Lanthanoids are chemically similar to each other, so that a diamagnetic reference can be obtained easily by substitution of the paramagnetic ion with Lu(III) or La(III). Furthermore, the ionic radius of Y(III) is practically the same as that of Ho(III), making it an excellent diamagnetic reference for the heavy lanthanoid ions ranging from Tb(III) to Yb(III). For this reason, the present article makes no formal distinction between Ln(III) and Y(III) ions. In the case of nitroxide radicals, the diamagnetic reference is usually obtained by chemical reduction to the corresponding hydroxylamine, which is readily achieved with ascorbic acid. Alternatively, diamagnetic control probes (such as hydroxylamine derivatives) can be used that are chemically similar to the nitroxide probes.⁶⁵ As the properties of 3d block ions vary more than those of Ln(III) ions, it is more difficult to identify suitable diamagnetic references. Zn(II) is commonly used as a diamagnetic reference for Co(II) and Mn(II) because charge and size are similar, and Ga(III) has been successfully used as a diamagnetic reference for high-spin Fe(III).^{66,67}

1.3. Quantitative Description of Paramagnetic Effects in NMR Spectroscopy

Mathematical descriptions of the different paramagnetic effects are well established and enable their quantitative interpretation. Hyperfine shifts, such as PCSs, are reported as the change in chemical shift (measured in ppm) caused by the presence of a paramagnetic center. RDCs are measured in hertz, reporting on the change in multiplet splitting observed due to weak paramagnetic alignment of the molecules with the magnetic field. PREs and cross-correlated relaxation effects are measured in s $^{-1}$, presenting the paramagnetic enhancement in longitudinal (R_1) or transverse (R_2) relaxation rates of nuclear spins over a corresponding diamagnetic reference, with R_1 and R_2 being the inverse of the T_1 and T_2 relaxation times. The basic equations governing the various paramagnetic effects have been treated in numerous review articles and books.^{7–9,11,13,19,68} In the following, we present a summary to highlight the salient features together with correction terms that may become relevant in special circumstances.

1.3.1. Pseudocontact Shift (PCS). The PCS of a nuclear spin can be described by eq 1⁷

$$\Delta\delta^{\text{PCS}} = \frac{1}{12\pi r^3} \left[\Delta\chi_{\text{ax}} (3 \cos^2 \theta - 1) + \frac{3}{2} \Delta\chi_{\text{rh}} (\sin^2 \theta \cos 2\phi) \right] \quad (1)$$

$$\Delta\chi_{\text{ax}} = \chi_z - \frac{\chi_x + \chi_y}{2} \quad \text{and} \quad \Delta\chi_{\text{rh}} = \chi_x - \chi_y \quad (2)$$

where r is the distance between the paramagnetic center and the nuclear spin, and the angles θ and ϕ are the polar angles describing the location of the nucleus with respect to the principal axes of the χ tensor. The axial component of the magnetic susceptibility anisotropy, $\Delta\chi_{\text{ax}}$, and the rhombic component $\Delta\chi_{\text{rh}}$ are defined by eq 2, where χ_x , χ_y , and χ_z denote the values of magnetic susceptibility along the respective principal axes of the χ tensor (Figure 1A). Being defined as the anisotropic component of the χ tensor, the $\Delta\chi$ tensor is spanned by principal axes that are aligned with those of the χ tensor. Use of eq 1 for calculating the PCS of a nuclear spin in a given molecular structure requires the prior knowledge of the location of the paramagnetic center and the orientation of the $\Delta\chi$ tensor relative to the molecule. The orientation of the $\Delta\chi$ tensor is commonly described by three Euler angles (α , β , and γ) and, in the case of proteins, reported relative to coordinates deposited in the protein data bank (PDB). Together with the $\Delta\chi_{\text{ax}}$ and $\Delta\chi_{\text{rh}}$ values, eight parameters are thus required to define the $\Delta\chi$ tensor relative to a set of atomic coordinates. They can be obtained by using the experimentally measured PCS data of at least eight different nuclear spins to fit the $\Delta\chi$ tensor to the molecular structure. A number of software packages^{69–73} are available to perform this fit. In practice, good fits require at least three times more PCS data than the minimum of eight, in which case it is also possible to obtain a measure of the uncertainties of the $\Delta\chi$ tensor parameters by Monte Carlo random variation of the data input.

To fit $\Delta\chi$ tensors to molecular structures and calculate correction terms arising from cross-correlation effects and molecular alignment, it is most convenient to use a matrix representation of the χ tensor. The isotropic component of the χ tensor is given by

$$\chi_{\text{iso}} = \frac{\mu_0 g_e^2 \mu_B^2 [S(S+1)]}{3k_B T} \quad (3)$$

where μ_0 is the induction constant and g_e and S are the electronic g factor and spin, respectively; their values are replaced by g_j and J for lanthanoid ions.⁷⁴ μ_B is the Bohr magneton; k_B denotes the Boltzmann constant, and T is the absolute temperature. Including the anisotropic components of the χ tensor, a concise mathematical description of the dipolar shift tensor σ at a given position r and distance r from the paramagnetic center can be written as

$$\sigma = \frac{1}{4\pi} \left[3 \frac{r \otimes r^T}{r^5} - \frac{\mathbb{I}_3}{r^3} \right] \cdot \chi$$

$$= \frac{1}{4\pi r^5} \begin{bmatrix} (3x^2 - r^2) & 3xy & 3xz \\ 3xy & (3y^2 - r^2) & 3yz \\ 3xz & 3yz & (3z^2 - r^2) \end{bmatrix} \cdot \begin{bmatrix} \chi_{xx} & \chi_{xy} & \chi_{xz} \\ \chi_{xy} & \chi_{yy} & \chi_{yz} \\ \chi_{xz} & \chi_{yz} & \chi_{zz} \end{bmatrix} \quad (4)$$

where \mathbb{I}_3 denotes the 3×3 identity matrix, \otimes denotes the Kronecker product, and x , y , and z the coordinates of the nucleus relative to the origin of the χ tensor.^{7,73} (Note that the dipolar shift tensor is the negative of the dipolar shielding tensor.) The full χ tensor including its isotropic component is needed to calculate PREs, but the $\Delta\chi$ tensor suffices to calculate PCSs. The $\Delta\chi$ tensor in matrix representation is the traceless part of the χ tensor (i.e., the same as the χ tensor, except that each diagonal element is reduced by their average). In the representation of eq 4, the PCS is given by the trace of the shielding tensor

$$\Delta\delta^{\text{PCS}} = \frac{1}{3} \text{Tr}[\sigma] \quad (5)$$

which can also be expressed in terms of $\Delta\chi$ tensor elements

$$\Delta\delta^{\text{PCS}} = \frac{1}{4\pi r^5} [x^2 - z^2, y^2 - z^2, 2xy, 2xz, 2yz] \cdot \begin{bmatrix} \Delta\chi_{xx} \\ \Delta\chi_{yy} \\ \Delta\chi_{xz} \\ \Delta\chi_{yz} \end{bmatrix} \quad (6)$$

Eq 6 illustrates how separating the position coordinates of the PCS equation (eq 5) from the $\Delta\chi$ tensor leads to a linear form of the equation with 5 unique elements in a column vector defining the tensor. Equation 6 thus allows $\Delta\chi$ tensor fits for a fixed position by singular value decomposition. Combined with the three coordinates of the metal center relative to the coordinates of the molecule, a $\Delta\chi$ tensor fit thus requires determining eight parameters.

1.3.2. Residual Dipolar Coupling (RDC). In isotropic solutions, internuclear dipolar couplings average to zero due to fast molecular tumbling, but RDCs re-emerge upon partial alignment of the molecules in a magnetic field. RDCs are most easily measured for one-bond scalar couplings, where they manifest in altered distance (measured in Hz) between well-separated multiplet components.²⁷ If the molecular alignment is caused by a paramagnetic center with anisotropic magnetic susceptibility, the alignment tensor A is directly proportional to the $\Delta\chi$ tensor, with the same axes directions⁷

$$A = \frac{B_0^2 \Delta\chi}{15\mu_0 k_B T} \quad (7)$$

where B_0 is the magnetic field strength. The residual dipolar coupling between two nuclear spins I and K is described by^{7,75}

$$D_{IK} = -\frac{hB_0^2\gamma_I\gamma_K}{240\pi^3 r_{IK}^3 k_B T} \left[\Delta\chi_{ax} (3 \cos^2 \Theta - 1) + \frac{3}{2} \Delta\chi_{th} \sin^2 \Theta \cos 2\Phi \right] \quad (8)$$

where γ_I and γ_K denote the gyromagnetic ratios of the nuclear spins I and K , respectively, r_{IK} is the internuclear distance, h Planck's constant, and the angles Θ and Φ determine the orientation of the I – K vector relative to the $\Delta\chi$ tensor (Figure 1B). The RDC, thus, yields information about the orientation of the internuclear vector in the frame of the $\Delta\chi$ tensor independent of the distance of the nuclei from the paramagnetic center. RDCs allow fitting the alignment tensor to a protein structure and, once the tensor frame has been determined, the RDCs can be used to establish bond vector orientations within a protein structure. As reorientational motions of a bond vector reduce the RDC by averaging over different orientations, RDC measurements can be used to study protein dynamics.^{27,76}

1.3.3. Residual anisotropic chemical shift (RACS). The weak molecular alignment in the magnetic field caused by anisotropic magnetic susceptibilities not only produces RDCs, but also changes the chemical shifts of nuclear spins that feature significant CSA tensor anisotropies by rendering the averaging over different molecular orientations incomplete. The change in chemical shift caused by such RACS effects poses a limit on the accuracy with which PCSs can be measured. If the structure of the molecule is known, the CSA tensors associated with individual nuclei can be taken into account by estimating a RACS correction for each PCS measured. The corrections can be significant at high magnetic field strength,⁷⁷ as the degree of molecular alignment depends on the square of the magnetic field (eq 7).

1.3.4. Residual Anisotropic Dipolar Shift (RADS). Residual anisotropic dipolar shifts present another effect by which experimentally measured PCS data may have to be corrected to obtain the pure PCS described by eq 1. Like the RACS effect, the RADS effect arises from weak molecular alignment in the magnetic field, which leads to incomplete averaging of anisotropic chemical shifts. Paramagnetic centers with a Curie spin create a dipolar shielding tensor at the site of the nuclear spin, which is generally anisotropic and similar to a CSA tensor. As molecular alignment results in nonuniform averaging of the dipolar shielding, the PCS produced by the effective DSA tensor is no longer fully represented by eq 1. Fortunately, RADS corrections can be calculated and, importantly, are barely measurable in practice.⁷

1.3.5. Saturation Effect. Finally, saturation of magnetic moments at high magnetic field strength renders the $\Delta\chi$ tensor and, hence PCSs, field dependent. This effect is small even at 18.8 T (800 MHz for ¹H NMR) but may become more important at much higher field strengths.⁷⁸

1.3.6. Paramagnetic Relaxation Enhancement (PRE). Spin–spin interactions are an important source of nuclear relaxation. Due to the large magnetic moment of unpaired electrons, electron–nuclear spin interactions easily provide a dominant source of nuclear relaxation. The SBM relaxation mechanism^{59,79,80} applies to all paramagnetic centers. For paramagnetic centers with unpaired electrons that relax rapidly within the rotational correlation time of the molecule, the Curie spin mechanism^{60,61} provides an additional and often dominant source of PREs. SBM relaxation describes the effect from dipole–dipole coupling between electron and nuclear spins, as it

drives nuclear relaxation by the variation in dipolar fields due to molecular reorientation and changes in the electronic spin state due to longitudinal electron relaxation. The paramagnetic enhancements in longitudinal (R_1^{SBM}) and transverse (R_2^{SBM}) relaxation rates can be described by

$$R_1^{\text{SBM}} = \frac{2}{15} \left(\frac{\mu_0}{4\pi} \right)^2 \frac{\gamma_I^2 g_e^2 \mu_B^2 S(S+1)}{r_{IS}^6} \left(\frac{3\tau_c}{1 + \omega_I^2 \tau_c^2} + \frac{7\tau_c}{1 + \omega_S^2 \tau_c^2} \right) \quad (9)$$

$$R_2^{\text{SBM}} = \frac{1}{15} \left(\frac{\mu_0}{4\pi} \right)^2 \frac{\gamma_I^2 g_e^2 \mu_B^2 S(S+1)}{r_{IS}^6} \left(4\tau_c + \frac{3\tau_c}{1 + \omega_I^2 \tau_c^2} + \frac{13\tau_c}{1 + \omega_S^2 \tau_c^2} \right) \quad (10)$$

$$\tau_c^{-1} = \tau_r^{-1} + \tau_s^{-1} \quad (11)$$

where r_{IS} is the distance between nuclear and electron spin and ω_I and ω_S are the nuclear and electron Larmor frequencies, respectively. Electronic spin flips are accounted for by the effective correlation time τ_c , which is composed of the rotational correlation time τ_r and the electronic lifetime τ_s as shown by eq 11. Terms in eqs 9 and 10 that depend on the electron frequencies can usually be neglected unless the lifetime of the electronic spin states is sufficiently short to approach the inverse of the electron precession frequency. The steep distance dependence of the PRE makes it very sensitive to changes in r_{IS} , which has been exploited to detect little populated conformations and protein–ligand complexes in solution.^{81–83}

Curie spin relaxation likewise results from dipole–dipole coupling between electron and nuclear spins, but it arises from the dipolar field created by the average electronic polarization due to higher populations of lower-energy as opposed to higher-energy electronic spin states.⁸⁴ Curie spin relaxation applies to systems in which electron relaxation occurs in a time short compared with the rotational correlation time of the molecule. Nuclear relaxation is insensitive to magnetic fields fluctuating at rates that are orders of magnitude faster than the Larmor frequency, such as the very fast electronic spin flips associated with Curie spins. Curie spin relaxation is driven by rotational tumbling, with the rotational correlation time τ_r characterizing the time-dependent modulation. Because of a quadratic dependence on the nuclear Larmor frequency, Curie spin relaxation can outweigh SBM relaxation for high-molecular weight systems in a strong magnetic field^{85,86} (note that the calculations of Table 2 in ref 85 were performed for $\tau_r = 10$ ns instead of $\tau_r = 1$ ns as stated in the original) while decreasing noticeably with increasing temperature.⁹ A quantitative description of PREs by Curie spin relaxation is given by eqs 12 and 13 for longitudinal (R_1^{Curie}) and transverse (R_2^{Curie}) relaxation rates.

$$R_1^{\text{Curie}} = \frac{2}{45} \left(\frac{\mu_0}{4\pi} \right)^2 \frac{\omega_I^2 g_e^4 \mu_B^4 S^2(S+1)^2}{(k_B T)^2 r_{IS}^6} \left(\frac{3\tau_r}{1 + \omega_I^2 \tau_r^2} \right) \quad (12)$$

$$R_2^{\text{Curie}} = \frac{1}{45} \left(\frac{\mu_0}{4\pi} \right)^2 \frac{\omega_I^2 g_e^4 \mu_B^4 S^2(S+1)^2}{(k_B T)^2 r_{IS}^6} \left(4\tau_r + \frac{3\tau_r}{1 + \omega_I^2 \tau_r^2} \right) \quad (13)$$

Eqs 9–13 describe the situation of isotropic magnetic moments associated with the paramagnetic center. In general, however, magnetic moments often are anisotropic. At the atomic level, such anisotropies usually arise when the magnetic moment of a paramagnetic metal ion depends not only on the electronic spin but also on spin–orbit couplings and therefore on the ligand field. This situation is particularly prominent for paramagnetic lanthanoid ions (except Gd(III)) and therefore their time-averaged magnetic moment is best described by an anisotropic molecular magnetic susceptibility tensor. For complexes with Sm(III) and Eu(III) ions, the situation is further complicated by a significant population of low-lying excited electronic states with different paramagnetic characteristics, which is manifested by a temperature dependence that is less steep than predicted by eqs 12 and 13.⁷⁴

An extension to the SBM theory has recently been described, which accounts for anisotropic dipolar spectral density in terms of a spectral power density tensor.⁸⁷ As this tensor usually cannot be derived theoretically, however, this extended theory depends on a greater number of parameters to be fitted to the experimental data. The anisotropic SBM theory is relevant for SBM relaxation generated by lanthanoid ions with anisotropic χ tensors. For these ions, however, Curie spin relaxation often exceeds SBM relaxation, making it difficult to detect the anisotropy effects in the SBM relaxation contribution.

The impact of anisotropic χ tensors on Curie spin relaxation can be calculated.⁶¹ Using the matrix formalism of eq 4, the R_1^{Curie} and R_2^{Curie} rates can be written

$$\Lambda^2 = (\sigma_{xy} - \sigma_{yx})^2 + (\sigma_{xz} - \sigma_{zx})^2 + (\sigma_{yz} - \sigma_{zy})^2 \quad (14)$$

$$\begin{aligned} \Delta^2 = & \sigma_{xx}^2 + \sigma_{yy}^2 + \sigma_{zz}^2 - \sigma_{xx}\sigma_{yy} - \sigma_{xx}\sigma_{zz} - \sigma_{yy}\sigma_{zz} \\ & + \frac{3}{4}[(\sigma_{xy} + \sigma_{yx})^2 + (\sigma_{xz} + \sigma_{zx})^2 + (\sigma_{yz} + \sigma_{zy})^2] \end{aligned} \quad (15)$$

$$R_1^{\text{Curie}} = \frac{1}{2}\Lambda^2\omega^2 \left[\frac{\tau_r}{1 + 9\tau_r^2\omega^2} \right] + \frac{2}{15}\Delta^2\omega^2 \left[\frac{\tau_r}{1 + \omega^2\tau_r^2} \right] \quad (16)$$

$$R_2^{\text{Curie}} = \frac{1}{4}\Lambda^2\omega^2 \left[\frac{\tau_r}{1 + 9\tau_r^2\omega^2} \right] + \frac{1}{45}\Delta^2\omega^2 \left[4\tau_r \frac{3\tau_r}{1 + \omega^2\tau_r^2} \right] \quad (17)$$

where ω refers to the Larmor frequency of the nuclear spin. The anisotropy effects tend to be small and not usually apparent in PRE measurements. Limited evidence has been found in cross-correlation effects.⁸⁸

1.3.7. Cross-correlated Relaxation (CCR). **1.3.7.1. DSA/CSA Cross-correlation.** For systems with Curie spins, the dipolar field emanating from the paramagnetic center contributes a variable magnetic field at the site of the nuclear spin, which drives nuclear relaxation in a way akin to chemical shift anisotropy. This analogy is highlighted by describing Curie spin relaxation as DSA relaxation. The matrix formalism of eq 4 automatically includes the effect from cross-correlation between DSA and CSA relaxation, as the effective shielding tensor at the site of the nuclear spin, σ^{eff} , is the sum of the DSA and CSA tensors. Experimentally, the PRE including cross-correlated relaxation is obtained as usual by subtracting the relaxation rate measured for the diamagnetic reference, $R(\sigma^{\text{CSA}})$, from the relaxation rate in the paramagnetic state, $R(\sigma^{\text{eff}})$, where both terms can be calculated using eqs 14–17.

$$R^{\text{Curie-CSA}} = R(\sigma^{\text{eff}}) - R(\sigma^{\text{CSA}}) \quad (18)$$

For nuclei for which CSA relaxation is the main relaxation mechanism in the diamagnetic state, the contribution by the DSA/CSA cross-correlation effect can be dominant. In this case, it is possible that the σ^{eff} tensor becomes more isotropic than the CSA tensor because of fortuitous compensation by the DSA tensor, which is manifested by lesser relaxation rates in the paramagnetic than in the diamagnetic state. This was predicted in 2004⁸⁵ and experimentally observed for the first time in 2016.⁵⁸ The effect was demonstrated with negative PREs of ¹⁵N nuclei.

1.3.7.2. DSA/DD Cross-correlation. Just as CSA/DD cross-correlation effects give rise to differential broadening effects of individual multiplet components (which has been exploited, for example, in TROSY spectra), DSA/DD cross-correlation effects also create differential broadening effects of multiplet components.⁸⁹ Using the matrix representation of eq 4, the cross-correlation effect can be calculated for the example of a ¹H spin that is bonded to a ¹⁵N nucleus by

$$\sigma_N = \frac{1}{B_0} \frac{\mu_0}{4\pi} \gamma_N \hbar I \left[3 \frac{r_{\text{HN}} \otimes r_{\text{HN}}^T}{r_{\text{HN}}^5} - \frac{\mathbb{1}_3}{r_{\text{HN}}^3} \right] \quad (19)$$

$$\sigma_{\uparrow} = \sigma + \sigma_N \quad (20)$$

$$\sigma_{\downarrow} = \sigma - \sigma_N \quad (21)$$

$$R^{\text{Curie-DD}} = R^{\text{Curie}}(\sigma_{\uparrow}) - R^{\text{Curie}}(\sigma_{\downarrow}) \quad (22)$$

where σ_N denotes the shift tensor at the site of the ¹H spin, I , that originates from the dipolar field of the ¹⁵N spin and adds either positively or negatively to the full shielding tensor of the ¹H spin, depending on the spin state of the ¹⁵N nucleus (as described by σ_{\uparrow} and σ_{\downarrow}). With the complete shift tensors at hand, the differential line broadening observed between the doublet components of the ¹H spin, $R^{\text{Curie-DD}}$, is readily calculated by using eqs 4, 14–17, and 19–22.

CSA/DD cross-correlation effects only appear in NMR spectra recorded without broadband decoupling. The value of the associated structural information has been demonstrated in a 3D structure of calculation of cytochrome *c'* from *Rhodobacter capsulatus* using paramagnetic restraints only.⁹⁰

1.3.7.3. Accuracy of $\Delta\chi$ Tensor Determination. The matrix representation of the $\Delta\chi$ tensor of eq 6 facilitates the fitting of $\Delta\chi$ tensors to the atomic coordinates of the macromolecular structure by a linear least-squares fit, using experimentally observed PCSs. The quality of the fit is commonly described by a quality factor Q , where a low value indicates a good fit:

$$Q = \sqrt{\frac{\sum_i \left[\left(\sum_m [a_i^{\text{exp}} - a_{m,i}^{\text{cal}}] \right)^2 \right]}{\sum_i \left[\left(\sum_m [a_i^{\text{exp}}] \right)^2 \right]}} \quad (23)$$

where a^{exp} and a^{cal} are the experimental and calculated PCSs values, respectively, the index m indicates ensemble averaging of spins that are common between different models of the molecular structure, and the index i is for summation over all spins of the molecule. Alternative Q factors have been proposed.^{91,92} The Q factor proposed by Bashir et al.⁹¹ uses sums of experimental and calculated values in the denominator of eq 23 and, therefore, tends to be 2 times smaller. This definition has the advantage that it does not bias for calculated values that are smaller over those that are larger than the

experimental values, which matters in the case of poor fits with large differences between observed and calculated data, as can be the case with PREs. For example, for $(a^{\text{exp}}, a^{\text{cal}})$ of (10, 20) $Q = 1$, but for (20, 10), $Q = 0.5$. The adjusted Q factor yields 0.33 in both cases. Importantly, Q factors are meaningful only, if the number of fitted data greatly exceeds the number of variables, as very small values in the denominator can render its calculation unstable. Therefore, fitting algorithms usually attempt to minimize the root-mean-square deviation (RMSD) between calculated and experimental values rather than Q factor.

In the case of paramagnetic metal probes attached to the target molecule via a long and flexible linker, the range of positions assumed by the metal ion relative to the target calls for an equal range of $\Delta\chi$ tensors to be fitted. Fitting multiple tensors, however, usually is not possible as each $\Delta\chi$ tensor determination requires at least eight PCSs to fit the tensor parameters and accurate fits require significantly more PCSs. The attempt to fit more than a single $\Delta\chi$ tensor to the limited data available would easily lead to problems with overfitting. The convention to fit single effective $\Delta\chi$ tensors has two consequences. First, the location of the paramagnetic center obtained by the fit does not correspond to its real position. In fact, it is quite possible that, due to temporary close proximity of the paramagnetic metal ion, the PCSs observed for a range of nuclear spins near the attachment site of the paramagnetic tag are larger than expected for a single immobile metal position. In this case, fitting of the data by a single effective $\Delta\chi$ tensor tends to increase the $\Delta\chi$ tensor and indicate a metal position that is further away from the surface of the target molecule than expected. This does not invalidate the use of effective $\Delta\chi$ tensors to back-calculate PCSs, but the predictive value of PCSs decreases with increasing distance from the nuclear spins, whose PCS data were used to fit the $\Delta\chi$ tensor.⁹³ Notably, exceptionally good correlations between experimental and back-calculated PCSs can be obtained even with tags attached via flexible linkers.⁹⁴

The difficulty to determine accurate $\Delta\chi$ tensors for flexible probes makes it difficult to assign the magnitudes of reported $\Delta\chi$ tensors to intrinsic probe characteristics. When a large number of PCSs has been used for the $\Delta\chi$ tensor fit, fits performed for rigid probes tend to produce low Q factors. An alternative way of assessing the flexibility of a paramagnetic probe is to compare the alignment tensor obtained by fitting RDCs with the $\Delta\chi$ tensor obtained by fitting PCSs, as both should be proportional to each other (eq 7). In practice, however, the alignment tensor is almost always smaller than that derived from PCSs because (i) RDCs are very sensitive to the accuracy of the molecular coordinates and (ii) molecules are not rigid and the orientations of the internuclear vectors determining the RDCs are averaged due to molecular dynamics as reflected by order parameters below one.

1.4. Paramagnetic NMR of Biomolecules in the Solid State

Solid-state NMR spectroscopy of biological macromolecules such as microcrystalline, fibrillar and membrane proteins is a growing area of interest largely driven by recent advancements in magic angle spinning (MAS) technology.^{95,96} Rotation speeds beyond 100 kHz have been shown to allow the acquisition of ^1H -detected multidimensional NMR spectra with good resolution and sensitivity without the need of protein perdeuteration.^{97,98} The first paramagnetic macromolecules to be investigated by solid-state NMR were metalloproteins, such as Co(II) substituted matrix metalloproteinase-12, for which PCSs were measured to characterize protein structure.^{99–101} Co(II)-

substituted superoxide dismutase was also investigated by PCSs and PREs with ^1H detected ultrafast MAS to determine molecular structure.^{102,103} Paramagnetic tags attached to biomolecules offer similarly useful long-range structural information in the solid state as in solution, but there are differences in the paramagnetic effects.^{53,52}

1.4.1. Paramagnetic NMR Effects in Static and Rotating Solids. An unordered paramagnetic compound in the solid state features molecular orientations distributed uniformly. As the dipolar shielding tensor arising from the paramagnetic center (see eq 4) renders the chemical shift for the nuclear spin dependent on the orientation of the molecule, the resulting powder pattern in the NMR spectrum reflects the principal axes of the σ tensor.

For the case of rotating solids, MAS of the paramagnetic species achieves coherent averaging of the σ tensor. At low spinning rates, this tends to split the powder pattern into spinning sidebands.¹⁰⁴ When the spinning rate is greater than the dipolar shielding term, however, only the isotropic component of the σ tensor, which includes the PCSs, remains observable in the NMR spectrum and can be described by eq 1.¹⁰⁵

While incoherent averaging of the σ tensor in solution NMR underpins the Curie spin relaxation mechanism as described in eqs 12–13, the coherent averaging by MAS effectively removes this pathway as a source of PREs in solid-state NMR.¹⁰⁶ Therefore, PREs of nuclear spins are unaffected by the Curie mechanism (except for nuclei in close proximity to the paramagnetic center, where the σ tensor term becomes larger than the MAS rate) and SBM relaxation (eqs 9–10) presents the dominant source of PREs in solids. SBM relaxation is governed by the incoherent electronic correlation time T_{1e} and independent of the MAS rate.⁵⁹

2. SOURCES OF PARAMAGNETISM

This section presents a brief overview of the most frequently used paramagnetic metal ions and nitroxide radicals.

2.1. 3d Block Transition Metal Ions

Transition metals in the 3d block of the periodic table can be in various oxidation states with different numbers of unpaired electrons occupying the d-orbitals.¹⁰⁷ Therefore, these cations are great candidates for generating various paramagnetic effects. They are frequently found in proteins. It is estimated that more than 25% of the known proteins contain one or more transition metal ions.¹⁰⁸ Paramagnetic NMR has long been applied to investigate the metal binding sites of metalloproteins.^{109–111} The first 3D structure determination of a metalloprotein was performed on a heme protein containing a low-spin Fe(III) ion.¹¹² Proteins containing many other paramagnetic transition metal ions have been studied, with iron, cobalt, manganese, copper, and nickel ions figuring most frequently. These ions are discussed hereafter.

Transition metal ion complexes of 3d block elements are stable with 4–6 electron donor sites. Depending on the number of coordination sites, transition metal ions feature mainly three types of coordination geometries, which are tetrahedral, square pyramidal, and octahedral (Figure 2). The magnetic properties of 3d block complexes can be strongly affected by the ligands, the environments, and the coordination geometries. Some general properties are shown in Table 1. The unpaired electrons usually are partially delocalized to the ligand orbitals, which affects the magnetic properties. Iron is a good example. Fe(III) and Fe(II)

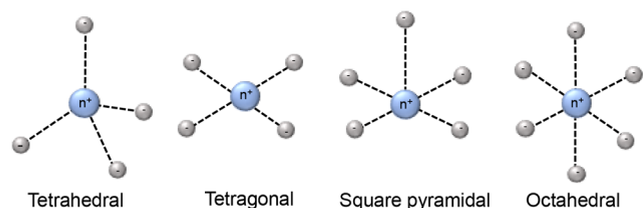


Figure 2. Schematic diagrams of 3d block ions coordination geometries.

are the most stable and commonly found oxidation states of iron. Both of them can be in paramagnetic high-spin states and Fe(III) is also paramagnetic in the low-spin state. With weak axial ligands, Fe(III) is in a high-spin state, whereas Fe(III) assumes a low-spin state with strong ligands. High-spin Fe(III) contains five unpaired electrons, and its electronic relaxation time can vary from 10^{-9} to 10^{-11} s.^{113,114} Therefore, it can generate sizable PCSs and PREs, depending on the ligands. The magnetic properties of low-spin state Fe(III) are quite different due to its short electronic relaxation time, which is below 10^{-11} s.^{113,114} As a result, the paramagnetism of low-spin Fe(III) is most prominently manifested in PCSs. Fe(II) has six electrons occupying d-orbitals, so when these electrons are paired, Fe(II) is diamagnetic. Its high-spin state has up to four unpaired electrons with a short electronic relaxation time (10^{-12} s), which causes PCSs.¹⁰⁷ Proteins containing iron ions have been well studied by paramagnetic NMR, for example cytochrome *c*, rubredoxin, and hemoglobin.⁶²

The electronic configuration of Co(II) is $3d^7$, which can be in a high-spin state containing three unpaired electrons or in a low-spin state containing a single unpaired electron. The electronic relaxation times of low-spin Co(II) are longer (10^{-9} – 10^{-10} s) than for the high-spin state ($\sim 10^{-12}$ s), thus low-spin Co(II) causes strong PREs and small PCSs.¹⁰⁷ In contrast, high-spin Co(II) can generate the largest PCSs among 3d block ions

combined with weak PREs. Mn(II) harbors one of its five unpaired electrons in each of the d-orbitals and has a long electronic relaxation time, causing the strongest PREs of all 3d block ions. Cu(II) is the most stable ionic state of copper. Three types of Cu(II) ions are found in proteins, which are distinguished by the Cu(II) ion coordination.^{15,62} In type-I copper(II) proteins, the ion is coordinated with trigonal (or distorted tetrahedral) geometry, involving at least one sulfur atom of cysteine and two nitrogens of two histidine residues.¹¹⁵ In type-II copper(II) proteins, the copper ion is coordinated by nitrogen and oxygen atoms with tetrahedral geometry. In the third type, two Cu(II) ions are antiferromagnetically coupled, decreasing the net magnetic susceptibility and electron relaxation time.¹¹⁶ Consequently, various paramagnetic effects were observed for Cu(II). Cu(II) contains one unpaired electron and generates small PCSs, but the PREs are strong, because of its relatively long electronic relaxation time (10^{-8} – 10^{-9} s). Therefore, Cu(II) is outstanding for PREs and it is widely used in EPR spectroscopy.^{107,117,118} The divalent ion of nickel has two unpaired electrons, generating small PCSs with sizable PREs.¹⁰⁷

2.2. Lanthanoid Ions

Lanthanoid ions were introduced into NMR studies of proteins decades ago, as agents for chemical shift changes and line broadening.^{119,120} The chemical properties of all Ln(III) ions are similar because the 4f-orbitals are shielded by 5s and 5p subshells, and their unpaired electrons do not participate substantially in different coordinating interactions with ligands. Ligands that bind Ln(III) ions thus have similar affinities for all lanthanoid ions. An overview of the paramagnetic properties of paramagnetic lanthanoid ions (excluding promethium, which is an unstable radioactive element) is presented in Table 2.

Gd(III) is unique because its seven unpaired electrons are distributed equally among the f-orbitals, producing a magnetic dipole moment that is independent of molecular orientation in

Table 1. Magnetic Properties of 3d Block and 4f Block Ions

ion	conf. ^a	J^b	τ_s^c	range ^d	PCSs ^e	PREs ^e	RDCs ^e
Fe(III) (HS)	[Ar]3d ⁵	5/2	10^{-9} – 10^{-13}	20	+	++	
Fe(III) (LS)	[Ar]3d ⁵	1/2	10^{-11} – 10^{-13}	15	++	+	
Mn(II)	[Ar]3d ⁵	5/2	10^{-8}	25		++	
Fe(II) (HS)	[Ar]3d ⁶	2	10^{-12}	15	+	+	
Co(II) (HS)	[Ar]3d ⁷	3/2	10^{-12}	25	++	+	
Ni(II)	[Ar]3d ⁸	1	10^{-10} – 10^{-12}	15	+	+	
Cu(II)	[Ar]3d ⁹	1/2	10^{-8} – 10^{-9}	20	+	++	
Ce(III)	[Xe]4f ¹	5/2	10^{-13}	10	+	+	
Pr(III)	[Xe]4f ²	4	10^{-13}	20	+	+	
Nd(III)	[Xe]4f ³	9/2	10^{-13}	10	+	+	
Sm(III)	[Xe]4f ⁵	5/2	10^{-13}	7	+	+	
Eu(III)	[Xe]4f ⁶	0	10^{-13}	15	+	+	
Gd(III)	[Xe]4f ⁷	7/2	10^{-8}	25		+++	
Tb(III)	[Xe]4f ⁸	6	10^{-13}	45	++++	+	+++
Dy(III)	[Xe]4f ⁹	15/2	10^{-13}	45	++++	+	+++
Ho(III)	[Xe]4f ¹⁰	8	10^{-13}	35	+++	+	++
Er(III)	[Xe]4f ¹¹	15/2	10^{-13}	30	++	+	+
Tm(III)	[Xe]4f ¹²	6	10^{-12} – 10^{-13}	50	+++	++	++
Yb(III)	[Xe]4f ¹³	7/2	10^{-13}	25	++	+	+
Nitr. ^f		1/2	10^{-7}	15		++	

^aElectronic configuration of the ions. ^bTotal angular momentum quantum number. ^cElectronic relaxation time (in seconds) reported in refs 29, 86, 735, and 736. ^dRange of paramagnetic effect (in Å) reported in ref 29, 107, and 86. ^e+ indicates the intensity of the effect based on refs 29, 107, and 86. No entry indicates not applicable or small effect. ^fNitr., nitroxide.

Table 2. Paramagnetic Properties of Paramagnetic and Nonradioactive Lanthanoid Ions^a

	Cerium	Praseodymium	Neodymium	Samarium	Europium	Gadolinium	Terbium	Dysprosium	Holmium	Erbium	Thulium	Ytterbium
	Ce	Pr	Nd	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Tb
χ_{iso}	5.6	11.2	11.4	0.6	~6	55.1	82.7	99.2	98.4	80.3	50.0	18.0
PRE												
PCS												
$\Delta\chi_{\text{ax}}$	2.1	3.4	1.7	0.2	2.4	0.0	42.1	34.7	18.5	12.2	26.0	8.5
$\Delta\chi_{\text{rh}}$	0.7	2.1	0.5	0.1	1.5	0.0	11.2	20.3	5.8	7.3	11.9	5.3
T_{1e}/ps	0.13	0.05	0.21	0.07	0.02	10 ⁵	0.25	0.24	0.21	0.19	0.27	0.16

^aThe radii of the yellow spheres correspond to the distance, where paramagnetic relaxation enhancement is predicted to broaden a ¹H NMR signal by 80 Hz on a 800 MHz NMR spectrometer, assuming a protein with a rotational correlation time of 15 ns and a temperature of 25 °C, calculated using eqs 10 and 13. The isotropic component of the χ tensors (χ_{iso} in 10⁻³² m³) were calculated using eq 3 with values for S and g_e taken from ref 74. The $\Delta\chi$ tensors reported for calbindin D_{9k}¹²⁵ are represented by isosurfaces of the PCSs drawn at ± 5 ppm. The values of the $\Delta\chi$ tensors are given in the unique tensor representation,⁶⁹ with their relative orientations corresponding to those reported for calbindin D_{9k}. The electronic longitudinal relaxation times were calculated for a magnetic field strength of 0.5 T following ref 737. They are not strongly field dependent.⁷³⁸ The T_{1e} time of gadolinium increases with the rotational correlation time and the square of the magnetic field strength.⁷³⁹ The scale bar at the left indicates a distance of 20 Å.

an external magnetic field. As its electronic relaxation time is long (>10⁻⁸ s) at field strengths >3 T,^{29,86} Gd(III) generates large PREs and enjoys increasing popularity in EPR as spin label.^{121–123} Although the remaining paramagnetic lanthanoid ions generate anisotropic χ tensors and produce all the paramagnetic effects discussed above, it has recently been shown that PRE measurements with Er(III), which generates large PREs associated with relatively small PCSs, can yield more accurate distance measurements than commonly used paramagnetic agents with long electronic relaxation times.¹²⁴ In general, Tb(III) and Dy(III) generate the largest $\Delta\chi$ tensors and, hence, cause the largest PCSs, while Tm(III) and Ho(III) produce medium-sized PCSs. Sizeable PCSs can also be observed with Yb(III) and Er(III). This general ordering, which follows theoretical expectations based on g_j factor and J quantum number⁷⁴ and has been experimentally verified for calbindin D_{9k} with one of the native calcium ions replaced by a Ln(III) ion,¹²⁵ is not always maintained as, for reasons not understood at present, some complexes of Tm(III) have been found to produce larger $\Delta\chi$ tensors than the same complexes with Dy(III) or Tb(III).^{126–129} Other lanthanoid ions are much less frequently used for paramagnetic NMR of proteins because their $\Delta\chi$ tensors are smaller.⁸⁶

2.3. Nitroxide Radicals

Nitroxide probes are organic molecules with five- or six-membered heterocyclic rings and a radical that is protected against homodimerization by bulky chemical groups (Figure 3). Because of their relatively small size combined with a long electronic relaxation time of a single unpaired electron and its reasonably well-defined localization, nitroxides are the most frequently used paramagnetic compounds in NMR to generate PRE distance restraints up to 20–25 Å.¹³⁰ The stability of the radical depends on the size of the ring and the substituents in the α position. In general, nitroxides in a five-membered ring are chemically more stable than in six-membered rings and bulkier

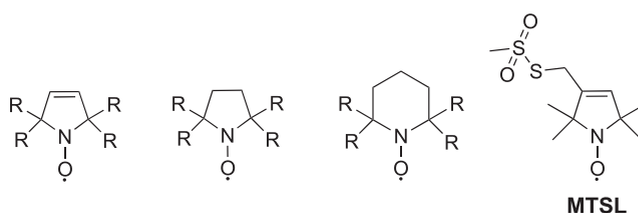


Figure 3. General structure of nitroxides. R represents an alkyl group.

groups in the α position promote stability by improved steric shielding. MTSL ((1-oxyl-2,2,5,5-tetramethyl-D-pyrroline-3-methyl)-methanethiosulfonate) is commercially available and the most popular nitroxide for protein labeling (Figure 3).

3. GENERAL OVERVIEW OF NATURAL AND CHEMICALLY GENERATED PARAMAGNETIC CENTERS

Paramagnetism of biomolecules can exist naturally or be introduced artificially by different strategies. Metalloproteins containing paramagnetic ions can readily be studied by NMR techniques tailored to paramagnetic samples. In some instances of diamagnetic metalloproteins, a diamagnetic metal ion can be replaced by a paramagnetic ion. Not all of the diamagnetic metalloproteins are suitable for metal ion substitution, however, and this approach does not work for nonmetalloproteins. Consequently, to gain access to the long-range structural restraints associated with paramagnetism, various methods have been devised to introduce paramagnetic centers. Two types of approaches can be distinguished. Paramagnetic centers can be added to the solution, resulting in solvent PREs. In this case, the PREs of protein nuclei are caused by nonspecific interactions with the paramagnetic relaxation agent in the solution. Such agents are designed to tumble freely and independently of the protein molecules, so that any anisotropy of the paramagnetic center will average to zero, thus yielding only PREs and

suppressing PCSs or RDCs. The probes for generating solvent-PRs are usually chemically synthesized.^{35,36}

Alternatively, a paramagnetic center can be attached covalently at a specific site on the protein. The two main methods of the covalent approach are the introduction of a genetically encoded metal binding site in the target protein and the chemical attachment of a paramagnetic center to the protein. Several principles need to be followed in the design of a suitable protein paramagnetic center. (i) The structure and properties of the target protein should be maintained. A highly charged or hydrophobic paramagnetic center as well as improper attachment sites can cause unfolding of the target protein and result in precipitation. Thus, small probes with low charge and high water solubility are preferred. (ii) To exploit the effects associated with paramagnetic metal ions featuring anisotropic χ tensors, the metal ion needs to be attached rigidly to the target protein, as the anisotropic effects decrease dramatically with the mobility of a paramagnetic center relative to the protein. Furthermore, movements of the paramagnetic center lead to averaging of the paramagnetic effects, which hampers the translation into structural information. (iii) The paramagnetic center should assume a single conformation. Many metal complexes show flexibility of parts of the coordinating cage, including exchange of coordinating groups. Different coordination states usually result in different orientations of the $\Delta\chi$ tensor and are thus prone to producing more than a single set of PCSs or RDCs (in the slow exchange limit) or line broadening (in the intermediate exchange regime). (iv) For probes that are linked via two identical tethers (e.g., to a pair of cysteine residues in the target protein), C_2 symmetry is important to avoid that attachment results in different isomers, each with its own $\Delta\chi$ tensor. (v) The metal affinity needs to be high and the probe needs to be stable and easily available, either commercially or by straightforward chemical synthesis.

3.1. Metalloproteins

3.1.1. Paramagnetic Metalloproteins. Among the metalloproteins with paramagnetic properties, iron and copper proteins are the best studied by paramagnetic NMR. Iron often occurs in either iron–sulfur clusters, such as in ferredoxin,^{113,131} or coordinated to heme rings, like in cytochromes.¹³² In FeS clusters, iron is bound to sulfurs from cysteine and inorganic sulfur.⁶² In heme, the macrocycle acts as a tetradentate ligand for the iron and provides space for additional axial ligands. Bertini and co-workers extensively studied iron proteins by paramagnetic NMR.^{112,133–139} The structure of oxidized *Saccharomyces cerevisiae* iso-1-cytochrome *c*, containing a low-spin Fe(III) ion, was the first paramagnetic metalloprotein for which the structure was refined with paramagnetic restraints.¹³⁹ The strategy used started with a known structure based on other restraints, such as NOEs, to fit the $\Delta\chi$ tensor of the paramagnetic center. This allowed using the PCSs as additional restraints in new rounds of structure calculations and refinement of the $\Delta\chi$ tensor based on the new structure. This iterative method has proven successful in yielding a more accurate 3D structure of the target protein.^{112,139}

3.1.2. Metalloproteins with Paramagnetic Substitution. In many cases, metals are either not paramagnetic or have inconvenient paramagnetic properties. In this situation, it may be possible to substitute the natural metal ion with another one characterized by different paramagnetic properties. For example, Cu(II) is usually ligated by histidine, cysteine, aspartic acid, or tyrosine residues, or a sulfide.^{115,140} As Cu(II) mainly generates

PREs, the NMR signals of the coordinating residues are very broad. By substitution of Cu(II) with Co(II), Donaire et al. studied coordination in the blue copper protein azurin¹⁴¹ and Bertini et al. investigated the metal binding site of stellacyanin.¹⁴² Also Ln(III) ions have been used for substitution of metal ions of similar ionic radius and coordination chemistry.¹⁴³ Calcium binding sites have been particularly successful in this respect and Ln(III) ions have frequently been successfully substituted for Ca(II) in the EF-hand motif.^{119,144} Most calcium proteins feature a pair of EF-hand motifs, but simultaneous substitution of both Ca(II) ions with two Ln(III) ions is unfavorable due to electrostatic repulsion. Therefore, calbindin D_{9k}, which possesses two Ca(II) binding sites in EF-hand motifs, is suitable for selective Ln(III) substitution into a single one of the Ca(II) binding sites.¹⁴⁵ Using calbindin D_{9k} as a model protein, the whole lanthanoid group was incorporated in this way, yielding a useful comparative study of their paramagnetic effects (Table 2).¹²⁵ Similarly, calmodulin, which harbors four calcium binding sites in four EF-hand motifs, has successfully been studied by paramagnetic NMR¹⁴⁶ and the Asn60Asp mutant was shown to promote the Ln(III) affinity and specificity of a specific Ca(II) binding site.^{145,147,148}

3.2. Genetically Encoded Metal Binding Sites

3.2.1. Natural Amino Acids or Peptides. Metal-generated paramagnetic structure restraints were initially explored in detail with paramagnetic metalloproteins and, having realized their exceptional value for 3D structure determinations of proteins, subsequently extended to diamagnetic metalloproteins by substitution with paramagnetic metal ions. In nonmetalloproteins, paramagnetic metal binding sites can be created by chemical modification of the target protein. Inspired by the strategy of substituting Ln(III) ions into EF-hands, lanthanoid-binding peptides (LBPs) have been proposed for adding a paramagnetic center to a protein (Table 3). Initially, a LBP with

Table 3. Frequently Used Lanthanoid Binding Peptides

lanthanoid binding peptide sequence	refs
DNDGDGKIGADE (“EF-hand”)	149
YIDTNNNDGWYEGDELLA	151, 152, 740–742
YIDTNNNDGAYEGDELLA	152
YIDTNNNDGWIEGDELLA	741, 742
YIDTNNNDGAYEGDELSSG	743
YIDTNNNDGWIEGDEL	741
GYIDTNNNDGWIEGDELY	154, 155
YVDTNNNDGAYEGDEL	157, 158, 626
GDYNKDGWYEELE	150

an EF-hand like motif was fused to the target protein at its N-terminus.¹⁴⁹ Subsequently, LBPs with improved lanthanoid ion binding affinity were identified by the Imperiali group and the number of residues reduced to 17.^{150–152} Double-lanthanoid-binding tags were shown to enable the binding of two Ln(III) ions simultaneously in a peptide with less than 40 residues.^{153,154} Because of the high mobility of terminal fusion tags, however, the paramagnetic effects in the target protein were greatly reduced by averaging. This situation was improved substantially by inserting an LBP into protein loops, as demonstrated for three different loops of interleukin-1 β (IL1 β) and confirmation of the tag structure by X-ray crystallography.¹⁵⁵ Coordinated with Gd(III) ions, these constructs can also be used for Gd–Gd distance measurement by EPR spectroscopy.¹⁵⁴ Disadvantages

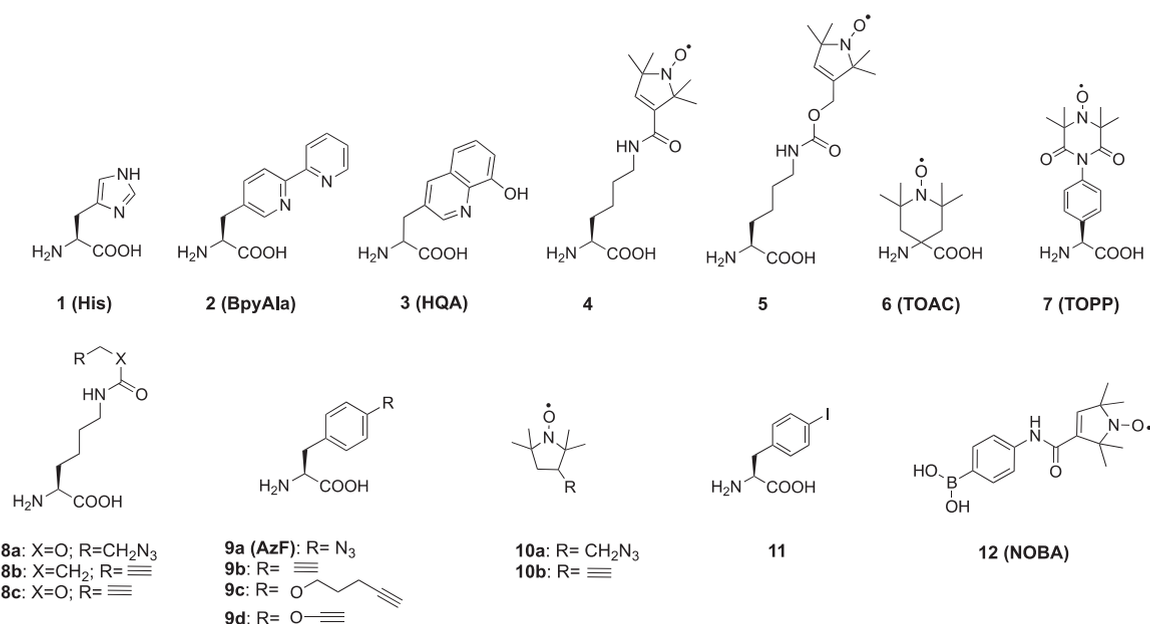


Figure 4. Chemical structures of some natural (1), noncanonical (2–9, 11) amino acids, and nitroxide radicals (10, 12).

of this approach are that it limits paramagnetic centers to termini or loop regions of the protein and any isotope labeling of the protein also labels the LBPs, which complicates the NMR spectra especially for the diamagnetic reference. In addition, this approach requires that the structure of the target protein is known, while the exact location of the lanthanoid ion is still difficult to predict in advance.

To overcome these drawbacks, LBPs were developed that can be attached by linking them to the protein chemically. Generally, these tags contain free thiol groups for attachment to cysteine residues.¹⁵⁶ They can be introduced anywhere by introducing a cysteine residue on the protein surface. Su et al. designed a series of LBPs with a cysteine residue for attachment to cysteine in the protein via a disulfide bond.¹⁵⁷ The $\Delta\chi$ tensor could be varied by including either a D- or L-cysteine in the LBP or varying the position of the cysteine residue in the LBP. Using the N-terminal domain of the *Escherichia coli* arginine repressor (ArgN) as model protein, large paramagnetic effects were obtained for all Ln(III) loaded LBPs with different tensor orientations for each of the tags,¹⁵⁷ but the mobility of the LBPs relative to the protein decreased the anisotropic paramagnetic effects similar to the LBP fusion method.¹⁵⁴ To reduce the mobility of the LBP tag, it has been proposed to anchor the tag at two sites.^{158,159} The first double-anchored LBP was designed by Saio et al. and combined a N-terminal fusion with a chemical linkage to the target protein via a disulfide bond.¹⁵⁸ Following fusion of the LBP to the N-terminus of the B1 immunoglobulin binding domain of protein G (GB1), a cysteine residue in the LBP was linked to a cysteine residue in GB1 by thiol activation using 5,5'-dithio-bis(2-nitrobenzoic acid) (DTNB) to form the second linkage. The double linkage was referred to as L2GB. As expected for a more rigid tag attachment, larger $\Delta\chi$ tensors were observed compared with the same LBP attached by fusion only (referred to as L1GB).¹⁵⁸

An alternative straightforward approach is to use the metal binding propensity of canonical amino acids like histidine, aspartic acid, and tyrosine to bind metal ions. For example, His₆ tags are routinely installed for protein purification and have been explored also for paramagnetic NMR. Unfortunately, like N-

terminal LBP fusions, this tag proved too mobile to yield good properties for paramagnetic NMR beyond generating PREs.¹⁶⁰ In contrast, dihistidine motives can generate better defined metal ion binding sites and have been used for EPR distance measurements between two Cu(II) ions.^{161–163} This approach was recently extended successfully to paramagnetic NMR studies, where a Co(II) ion bound to a dihistidine motif generated sizable PCSs in various proteins.¹⁶⁴ It was shown that dihistidine motives with good metal binding affinity can be installed either in α -helices or β -sheets.¹⁶⁴

3.2.2. Noncanonical Amino Acid and Their General Synthetic Approaches. Well over 100 noncanonical amino acids (ncAAs) with a wide variety of side chains have been incorporated into proteins by genetic encoding.^{165–167} Among these, some are capable of binding metal ions. For example, bipyridylalanine (**BpyAla**, Figure 4)¹⁶⁸ has successfully been used to endow proteins with new hydrolytic function or greater stability enabled by its capacity to bind divalent metal ions.^{169,170} Paramagnetic transition metal ions, such as Co(II), which prefer nitrogen ligands and require fewer coordination sites than Ln(III) ions, can be coordinated by **BpyAla**. A West Nile virus NS2B-NS3 protease (WNVpro) mutant containing a **BpyAla** residue in a loop was shown to generate PCSs following coordination with a Co(II) ion, but additional coordination was required from proximal side chains, as was demonstrated by mutation of nearby residues.¹⁷¹ 2-Amino-3-(8-hydroxyquinolin-3-yl)propanoic acid (**HQA**, Figure 4)¹⁷² has equally been explored as the basis for metal ion binding motives.^{173–178} Unfortunately, Ln(III) ions coordinated by **HQA** generally lead to quantitative protein precipitation,¹⁷⁸ but Mn(II) captured by **HQA** has successfully been used to generate PREs in membrane proteins.¹⁷³

A related approach uses site-specific incorporation of one or two phosphoserine residues, which is a naturally occurring amino acid produced by post-translational modification, but which can also be incorporated as a ncAA in response to an amber stop codon, owing to a recently developed genetic encoding system.^{179–181} A phosphoserine residue in conjunction with an aspartate or glutamate residue, or two

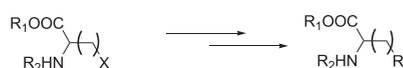
phosphoserine residues together with a glutamate residue can generate Ln(III) binding sites that position the metal ion very precisely on the target protein, generating substantial $\Delta\chi$ tensors.¹⁸² Unfortunately, the close proximity of charged amino acid side chains produces significant electrostatic repulsion, which can lead to unfolding of the protein and poor protein yields.

As discussed in section 2, not only paramagnetic metal ions but also nitroxide radicals constitute useful paramagnetic centers for generating PREs. Noncanonical amino acids containing a nitroxide radical have been synthesized (Figure 4, compounds 4–7)^{183–185} and some of them have been incorporated into proteins by genetic encoding.^{51,183} Alternatively, a nitroxide radical can be attached to a nCAA after incorporation in the target protein. In one method, an azido or alkynyl-bearing nCAA (Figure 4, 8a/9a or 8b/c/9b)^{186–188} is installed in the target protein and reacted with an alkynyl or azido-functionalized nitroxide radical (Figure 4, 10a or 10b),¹⁸⁸ using the copper-catalyzed azide–alkyne cycloaddition (CuAAC) reaction¹⁸⁹ to form a triazole linker. A similar strategy employs the Suzuki–Miyaura coupling reaction, where an iodide nCAA (Figure 4, 11) is reacted with a boronic acid spin label (NOBA, Figure 4, 12).¹⁹⁰ For nCAAs containing a nitroxide group from the beginning, reduction of the radical by the reducing conditions inside *E. coli* cells limits the yield of successful incorporation. Among the nitroxide radical nCAAs mentioned above, only TOAC was used as a paramagnetic center for NMR studies, to generate PREs in the complex between a peptide derived from focal adhesion kinase (FAK) and Src homology 3 (SH3) domain of Src kinase.¹⁹¹ Others were only used for EPR studies.^{51,183,188,192–195}

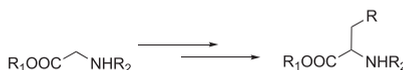
Two main strategies are followed in the synthesis of nCAA, which involve either the side chain modification of a natural AA or alkylation of a glycine equivalent (Scheme 1).^{196–198}

Scheme 1. Synthetic Strategies for Modification of Amino Acids

Method A: Modification of a natural AA



Method B: Alkylation of a glycine equivalent



Recently, additional strategies for nCAA synthesis have been published.^{199–201} Method A of Scheme 1 is most frequently applied to synthesize nCAA probes. *p*-Azido-*L*-phenylalanine, which is one of the most popular nCAAs, is synthesized by the first strategy with *L*-phenylalanine as the starting compound. Two approaches give the product in two steps (Scheme 2A).^{202–204} Similarly, compound 4 is synthesized by coupling reaction between *L*-asparagine and a nitroxide radical derivative (Scheme 2B).¹⁸³ BpyAla and HQA are nCAAs that can directly coordinate metal ions. Their synthetic routes differ from those mentioned above. Instead of starting from a natural AA, their synthesis starts from the compounds possessing the metal binding ability, which are modified with amido and carboxyl groups to form the final nCAA, as shown in Scheme 2C and D.^{168,172,205}

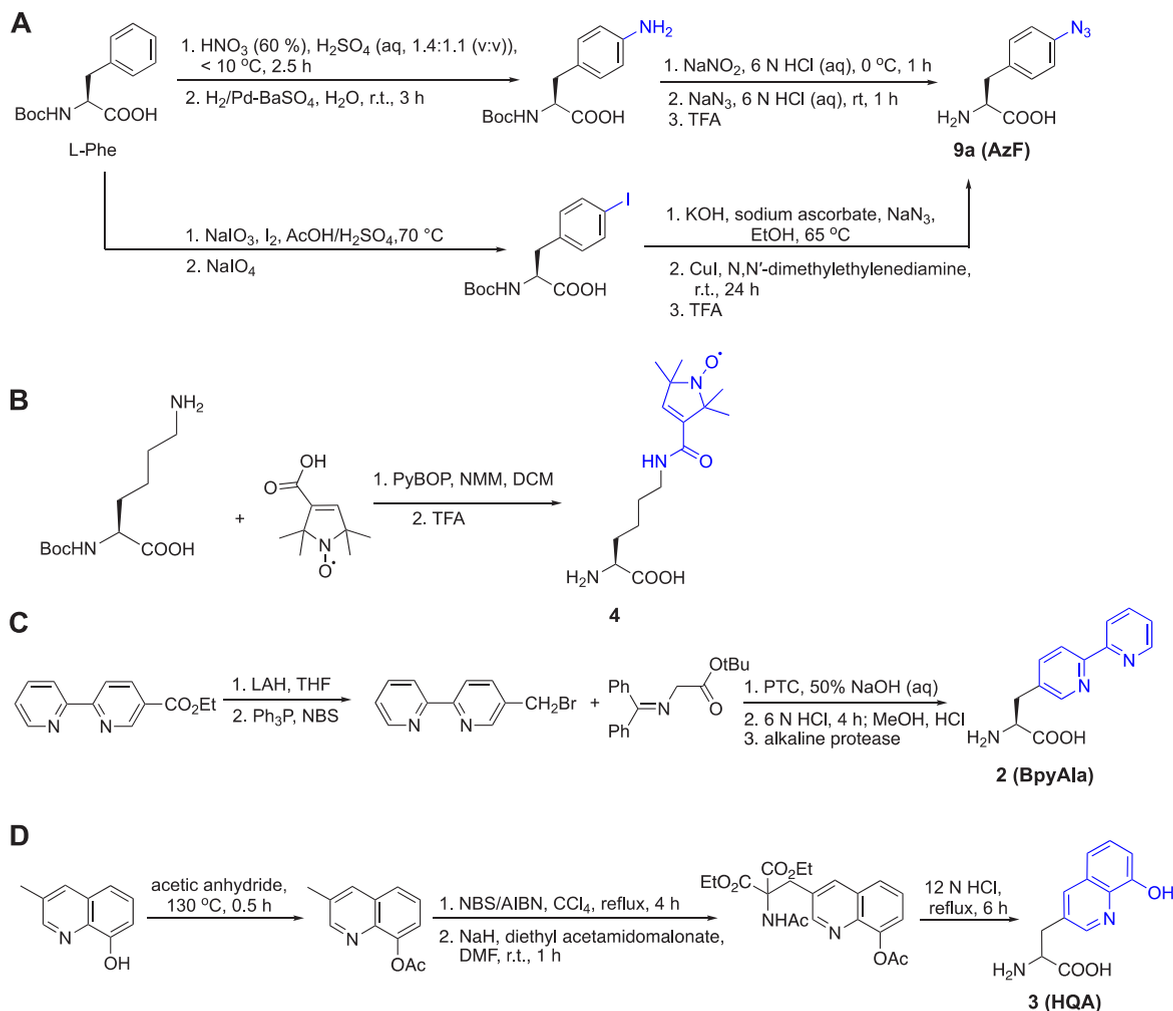
3.3. Synthetic Tags for Proteins

To obtain useful paramagnetic results, other than solvent PREs or RDCs, it is critical that the paramagnetic center is at a fixed distance and orientation relative to the nuclear spin and its molecular frame. Rapid distance and orientation variation lead to nonlinear averaging of the paramagnetic effects, making interpretation more complicated or impossible. Thus, the linkage between the paramagnetic center and the molecule needs to be as short and rigid as possible, contrary to, for example, the longer linkers that are often used in fluorescence spectroscopy. Second, it is critical that a paramagnetic center is attached to only a single, well-defined site on the molecule. The presence of more than one center greatly complicates the interpretation of PCS and PRE data. Similarly, if the tagging site is undetermined, the information content of PCS and PRE data becomes uncertain. As a consequence, tagging of proteins is best achieved on a residue with unique chemical reactivity on the accessible surface. For this reason, cysteine is by far the most popular residue type targeted for tagging, as many proteins lack surface exposed cysteine residues, so that unique sites can be engineered by site-directed mutagenesis. If exposed cysteines are present in the native protein, they can be replaced first by alanine or serine, usually without consequence for the structure or function of the protein. The general procedure requires reduction of any disulfide bridges using a reducing agent such as dithiothreitol, and removal of the reductant, before the tagging reaction can take place.

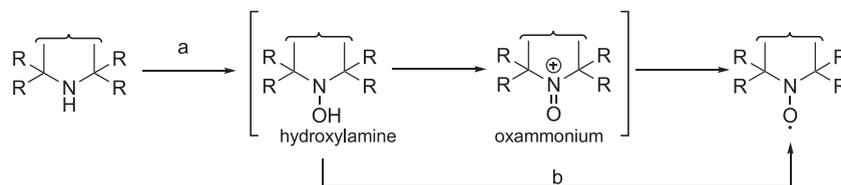
3.3.1. Nitroxide Probes. **3.3.1.1. Nitroxide Probes for Studies of Biomolecular Structure.** Nitroxides are extensively used as spin-labels for structural studies of proteins, protein–DNA complexes, and protein–RNA complexes by EPR and NMR spectroscopy. Two main factors need to be considered for the application of nitroxides in biological systems. The radical character must be stable with respect to the chemistry used to attach the tag to the target molecule. Nitroxides are stable under most nonreducing conditions but can nonetheless be prone to oxidation to an oxammonium cation or reduction to a hydroxylamine (see Scheme 3). Several ways have been reported to stabilize nitroxides.²⁰⁶ Besides increased chemical stability associated with five- versus six-membered heterocycles and bulky substituents adjacent to the nitroxide group, a nitroxide equipped with cycloalkanyl groups (Figure 5, 13) also tends to possess a longer electronic relaxation time than MTSL (Figure 3).²⁰⁷

The most efficient way of connecting a nitroxide label to a biomolecule is to use an activated thiol, such as methanethiosulfonate (MST)^{208,209} or a pyridylthio group,²¹⁰ which form disulfide bonds with cysteine residues. Disulfide bonds, however, are easily cleaved under reducing conditions. Therefore, maleimide^{211,212} and iodide^{212–214} nitroxide probes have been developed (Figure 5, 14–18) to form thioether bonds with cysteine residues, which are more stable under reducing conditions than disulfide bonds.

An important consideration is that any flexibility of the linker (Figure 6A) leads to averaging of the spin–spin interactions over a range of distances, reducing the accuracy of any distance measurements.^{215–218} Probe mobility can be restricted by adding a bulky chemical group on the heterocycle (Figure 5, 19),²¹⁹ introducing a second attachment group (Figure 5, 20–22),^{220–224} or incorporating a free radical containing unnatural amino acid into a sterically crowded site of the target protein (Figure 4, 4–7).^{183,195} In addition, nitroxides with alternative reactive functionalities have been reported, such as the azide

Scheme 2. Synthetic Routes of *p*-Azido-*L*-phenylalanine (A),^{202–204} Compound 4 (B),¹⁸³ BpyAla (C),²⁰⁵ and HQA (D)^{174,a}

^aThe functional groups of the ncAAs are colored in blue.

Scheme 3. Two Synthetic Approaches toward Nitroxide Radicals^a

^a(a) Oxidation with *m*-CPBA or H₂O₂/cat. Na₂WO₂. (b) Oxidation with MnO₂.

(Figure 5, 23–25)^{225,226} and alkyne (Figure 5, 26–30)^{227–229} containing probes amenable to “click” chemistry for attachment to, for example, *p*-azido-*L*-phenylalanine (AzF, Figure 4, 9a). Others contain an isothiocyanate (Figure 5, 31–32)^{230,231} or carbodiimide (Figure 5, 33)²³¹ group for reaction with a glutamate side chain, the hydroxylamine ether probe (Figure 5, 34)²³² to generate an oxime ether upon reaction with proteins containing a *p*-acetylphenylalanine residue, or a hydroxysuccinimide ester (Figure 5, 35)²⁰⁷ for reaction with a lysine side chain. These probes contain bulky groups in the vicinity of the nitroxide group and are well suited to minimize the movement of the radical relative to the protein frame.

3.3.1.2. General Synthetic Approaches toward Nitroxide Probes. Several methods have been established for the

preparation of nitroxides.²⁰⁶ In general, the nitroxide is formed by oxidation of a secondary amine with *meta*-chloroperbenzoic acid (*m*-CPBA) or H₂O₂ as oxidant (Scheme 3). The latter method requires a catalyst. In both methods, the amine group is first oxidized to form a hydroxylamine intermediate that is subsequently further oxidized to an oxammonium salt intermediate to finally yield the nitroxide. Alternatively, a mild oxidant, such as MnO₂, can be applied to oxidize a hydroxylamine to give the desired nitroxide (Scheme 3). The *m*-CPBA oxidation is fast because less polar solvents can be used and compounds with electron-deficient double bonds are tolerated under these conditions.²³³ In contrast, H₂O₂ in the presence of a catalyst (tungstate salt) requires the use of a polar solvent system, in which lipophilic substrates are poorly soluble.

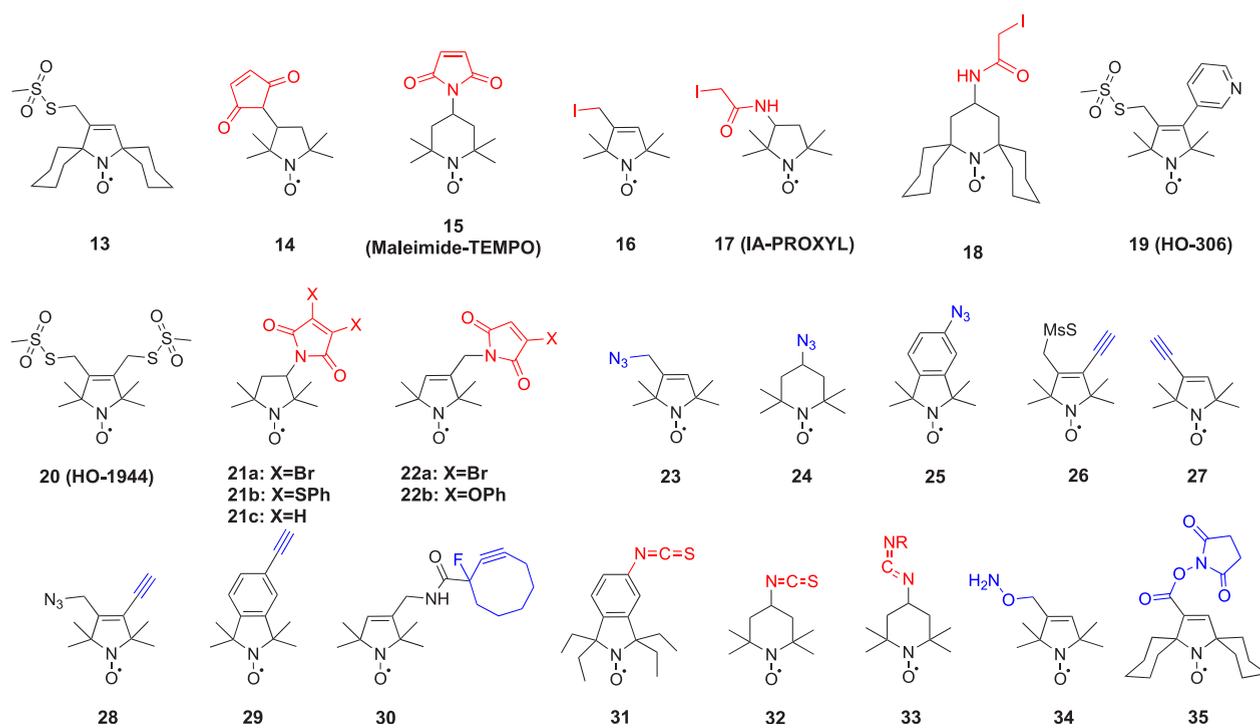


Figure 5. Structures of nitroxide probes. The functional groups for covalent attachment to cysteine via a thioether are highlighted in red and the functional groups for attachment to a nCAA are highlighted in blue. Probe 13²⁰⁷ is derived from MTS; 14²¹¹ and 15²¹² are maleimide functionalized nitroxides; probes 16,²¹³ 17,²¹³ and 18³⁹¹ are iodide functionalized nitroxides; probes 20,²²¹ 21,^{221,223} and 22²²⁴ are double-anchored nitroxides; probes 23,²²⁵ 24,²²⁵ and 25²²⁵ are azide functionalized nitroxides; probes 26,²²⁷ 27,²²⁷ 28,²²⁷ 29,²²⁸ and 30²²⁹ are alkyne functionalized nitroxides; probes 31²³⁰ and 32²³¹ are iso(thio)cyanide functionalized nitroxides; probe 33²³¹ is a carbodiimide functionalized nitroxide; probe 34²³² is a hydroxylamine ether functionalized nitroxide; probe 35²⁰⁷ is a nitroxide functionalized with a hydroxysuccinimide ester.

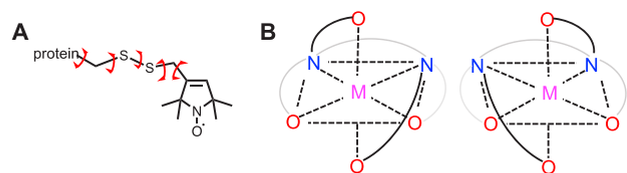


Figure 6. (A) Reaction of MTS with a cysteine residue of a protein generates a flexible linkage with five rotatable bonds (red arrows). (B) Schematic representation of two different enantiomeric EDTA complexes produced by different coordination of the metal ion. M denotes the metal ion. O denotes the carboxyl groups. Curved lines represent the ethylene groups and dashed lines trace the octahedral coordination.

As discussed in section 3.3.1, there are two factors that need to be considered for improving the stability of nitroxides, one being the ring size and the other the substituents in the α positions.^{206,234} In the following, we discuss the synthesis of different types of nitroxides with five- or six-membered rings and various substituents in the α -positions.

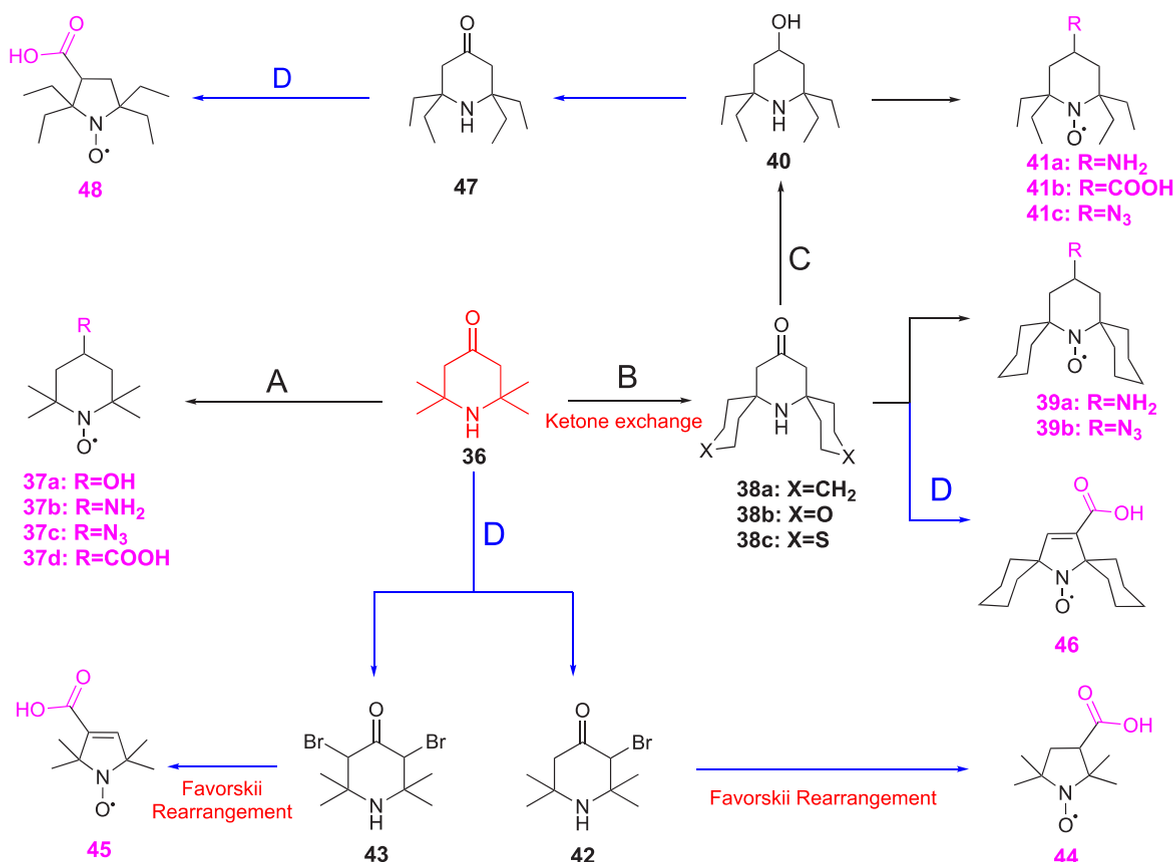
Tetramethylpiperidone (Scheme 4, 36), which is commercially available, is the most commonly used starting material for the synthesis of nitroxide probes, due to its reactive ketone group. It can readily be modified to provide water-soluble TEMPO derivatives (see Scheme 4) or embellished with a reactive functionality (Scheme 4, 37a–d).^{225,231,235} Sakai et al. successfully developed a mild method to exchange the tetramethyl groups of compound 36 by more bulky six-membered heterocyclic or homocyclic rings (Scheme 4, 38a–b).²³⁶ In this reaction ammonium chloride acts as catalyst and, by using ¹⁵NH₄Cl, the authors were able to study the mechanism

of this interesting exchange reaction that appears to involve two cross-aldol reactions, two fragmentations and two Michael additions, one of which acts as the final cyclization step.²³⁶ Similarly, the ketone group of compound 38a was transformed to produce compounds 39a–b (Scheme 4).²²⁵ The sulfur–carbon bonds of compound 38c can be removed by Raney nickel to give a tetraethyl substituted derivative (Scheme 4, 40).²³⁷ After oxidation of the NH of compound 40 to a nitroxide, the hydroxyl group can be converted to different functional groups (compounds 41a–c).^{225,238,239}

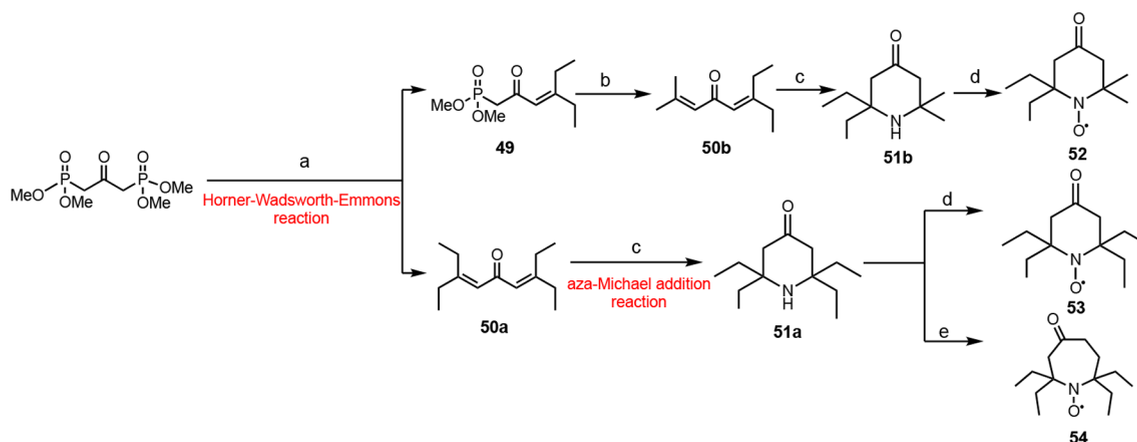
Tetramethylpiperidone is also a precursor for the synthesis of five-membered ring nitroxides via a Favorskii ring contraction.^{225,240} Tetramethylpiperidone 36 is first brominated to yield compound 42 or 43. Under basic conditions, compounds 42 and 43 give the pyrrolidine derivative 44 and the pyrroline 45, respectively (Scheme 4). A similar sequence of steps leads to compounds 46 and 48 (Scheme 4).^{225,233,237}

Another method based on the Horner–Wadsworth–Emmons reaction for making α -tetrasubstituted piperidones from a bisphosphonate has been reported (Scheme 5).²⁴¹ In this procedure, a monoeneone (Scheme 5, compound 49) and a dienone intermediate (Scheme 5, compound 50a) were generated. The monoeneone was further reacted to give nonsymmetric dienone (Scheme 5, compound 50b). Both dienones (Scheme 5, compounds 50a and 50b) further reacted with hydroxylamine through a double aza-Michael addition reaction to yield the corresponding tetra-substituted piperidones 51a and 51b, which were oxidized to give the nitroxides (Scheme 5, compounds 52–54).

Isoindolyl nitroxides are of special interest due to their benzannulated scaffold with diminished flexibility as compared

Scheme 4. Preparation of Various Nitroxides from Tetramethylpiperidone^a

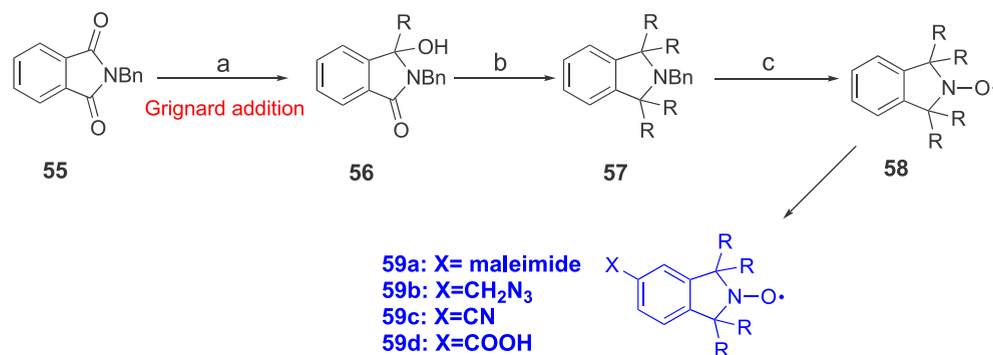
^a(A) i. NaBH₄, H₂O₂/cat. Na₂WO₂ (to give 37a);²²⁵ ii. reductive amination with NaBH₃CN (to give 37b);²³¹ iii. active amine with methanesulfonyl to substitute with NaN₃ (to give 37c);²²⁵ iv. carboxylation with tosylmethylisocyanide (to give 37d).²³⁵ (B) Cyclohexanone, NH₄Cl, H₂O₂/cat. Na₂WO₂ (to give 38a–c).²³⁶ (C) Raney–nickel (to give 40).²³⁷ (D) i. Bromination (to give 42, 43); ii. Favorskii rearrangement (to give 44, 45).^{225,240} Blue colored arrows indicate the synthetic route to five-membered ring nitroxides.

Scheme 5. Synthetic Route to α -Tetrasubstituted Piperidone from a Bisphosphonate Precursor^a

^a(a) i. LDA (lithium diisopropylamide) at 0 °C. ii. BuLi (butyllithium) at –35 °C. iii. excess 3-pentanone. (b) i. LDA, THF, ii. acetone. (c) i. NH₄OH at 105 °C. ii. *m*-CPBA. (d) Na₂WO₄, H₂O₂. (e) i. TMSCHN₂ ((diazomethyl)trimethylsilane), BF₃·OEt₂; ii. Na₂WO₄, H₂O₂.

to aliphatic nitroxide derivatives.²²⁵ *N*-benzylphthalimide (Scheme 6, compound 55) is the most popular starting material for the synthesis of isoindolinylnitroxides reported in the early 1980s, via the use of excess Grignard reagent to give 57.²⁴² A more efficient procedure than a quadruple Grignard addition was found later, which proceeded via isolation of the

monoaddition product (compound 56) and subsequent conversion into compound 57 (Scheme 6).²⁴³ The further transformation of the benzylamine to a nitroxide was achieved by debenzoylation and oxidation (Scheme 6, 58).^{244–246} The latter compound can readily be transformed into a probe, such as compound 59a/b (Scheme 6), which can be attached to a thiol

Scheme 6. Synthetic Routes Towards Isoindolinyl Nitroxide^a

^a(a) RMgBr. (b) Excess RMgBr. (c) i. Pd–C/H₂; ii. *m*-CPBA.

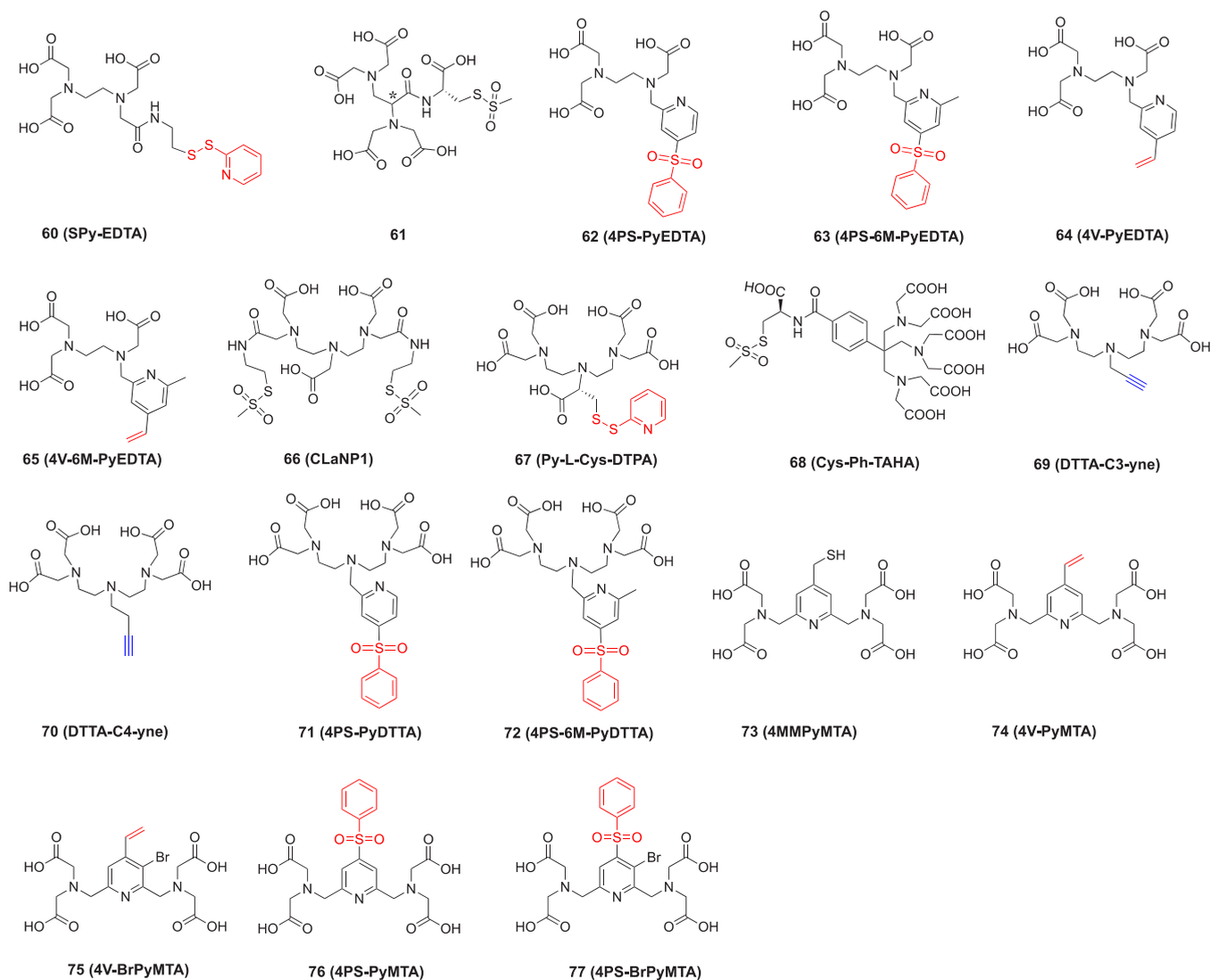
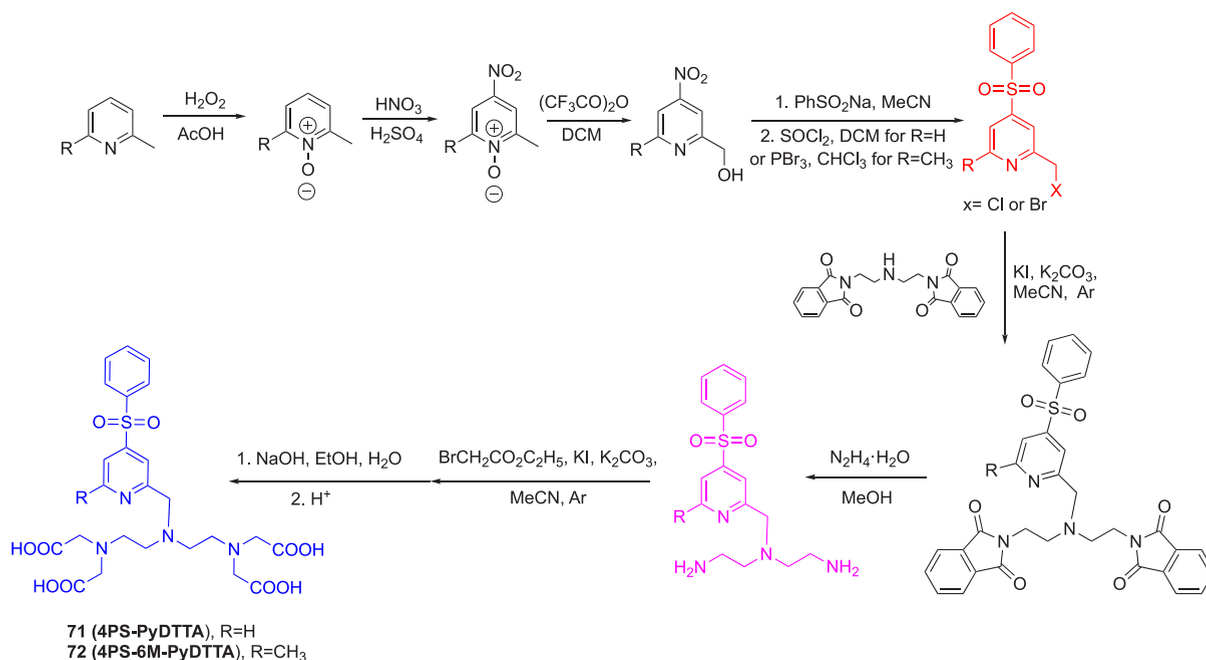


Figure 7. Structures of aminopoly(carboxylic acid) based probes. Functional groups for covalent attachment to a cysteine residue in the target protein via thioether formation are highlighted in red. Alkyne groups for attachment to an azide group in a ncAA via “click” reaction are highlighted in blue. Probes 60–65^{249–253} are EDTA based. Probe 61 has two chiral centers. Probe 66²⁵⁵ is double-anchored. Probes 73–77^{263–266} are PyMTA based. The asterisk identifies one of the chiral carbon atoms.

group of a biological macromolecule.²⁴⁴ Several other routes toward functionalized isoindolinyl nitroxides have been reported. One route involves bromination of the aromatic ring and removal of the benzyl protection group of compound 55,

followed by substitution of the bromide by a cyano or carboxyl group (Scheme 6, 59c/d).²⁴⁷ Another route involves the construction of the functionalized aromatic ring via a Diels–Alder reaction on a diene nitroxide²²¹ or following a higher

Scheme 7. Synthetic Route to Probes 71 (4PS-PyDTTA) and 72 (4PS-6M-PyDTTA).²⁶⁰

yielding stepwise procedure using a disubstituted pyrrole nitroxide as starting material, leading to compound **59d**.²⁴⁷

3.3.2. Aminopoly(Carboxylic Acid)-Based Probes.

Aminopoly(carboxylic acid)s have the ability to bind metals strongly and they can easily be modified with functional groups to enable linkage to biomolecules. In this section, the most commonly applied aminopoly(carboxylic acid) based probes and their synthetic strategies are discussed.

3.3.2.1. EDTA-Based Probes. EDTA (ethylenediaminetetraacetic acid) is a widely used chelating agent for a broad range of metal ions, due to its high metal ion binding affinity with four carboxylic acids and two amine groups. Its complexes with paramagnetic ions can be linked to proteins via a disulfide bond by functionalizing them with MTS or pyridylthio groups. The first EDTA based paramagnetic probe for proteins, **SPy-EDTA** (Figure 7, **60**), was reported by Ebright et al. in 1992.²⁴⁸ This probe soon became commercially available and its various metal ion complexes were used for protein paramagnetic NMR studies.^{249–251} These studies revealed two sets of PCSs generated by probe **60** loaded with Co(II) or Dy(III), as the metal complex produces two enantiomeric forms, which leads to diastereomers after linkage with the protein (Figure 6B).^{249,251} Subsequent introduction of chiral centers and a rigid attachment group to Ln(III)-EDTA resulted in single sets of PCSs, but the $\Delta\chi$ tensor was relatively small (Figure 7, **61**).^{251,252} Diastereomer formation is not a critical problem for DEER distance measurements and Su et al. (2015) designed EDTA based EPR probes with short linkers as Mn(II) tags (Figure 7, **62–65**), which yielded a narrow distribution of distances.²⁵³

3.3.2.2. DTPA-, DTTA-, and TAHA-Based Probes. Diethylenetriaminepentaacetic acid (DTPA) has a structure similar to EDTA, but features a higher metal binding affinity.²⁵⁴ DTPA functionalized with two MST groups, **CLaNP-1** (caged lanthanoid NMR probe 1, Figure 7, **66**), was designed as an NMR probe for proteins, using Yb(III) and Y(III) as paramagnetic and diamagnetic metal ions, respectively.²⁵⁵ **CLaNP-1** loaded with Yb(III) generated large PCSs, but multiple sets of PCSs were observed for each nuclear spin

observed in the protein, because the coordination cage assumes multiple conformations.²⁵⁵ DTPA coordinates lanthanoids in an octadentate fashion and forms a pair of enantiomers.²⁵⁶ In 2020, Su and co-workers showed that a combination of DTPA and L-cysteine (**Py-L-Cys-DTPA** (Figure 7, **67**))²⁵⁷ yields an open chain lanthanoid chelator that shows no exchange between conformations on the low millisecond time. Upon attachment to various sites on ubiquitin, large PCS and RDC were obtained, which were dependent on the site of attachment, likely due to a different degree of mobility for this single-arm tag. The probe **Cys-Ph-TAHA** (Figure 7, **68**)²⁵⁸ was based on C₃-symmetric TAHA (triaminohexaacetate). The paramagnetic properties of lanthanoid loaded **Cys-Ph-TAHA** were determined by using two ubiquitin mutants (T12C and S57C). PCSs and RDCs were observed. Two minor species were observed for **Cys-Ph-TAHA** labeled ubiquitin T12C, but not for ubiquitin S57C.²⁵⁸

Two DTTA (diethylene-triamine-tetraacetate) based protein probes (**DTTA-C3-yne** and **DTTA-C4-yne**, Figure 7, **69** and **70**)²⁵⁹ were reported, which possess an alkynyl group for “click” ligation to a genetically encoded nCAA containing an azide group, with formation of a triazole linker (Figure 4, **9a**). Tb(III) or Tm(III) loaded probes conjugated to ubiquitin mutants generated single sets of PCSs with a good correlation between experimental and back-calculated values. The triazole ring in the linker limits the number of rotatable bonds as it can form a conjugated double-bond system with the phenyl ring of the noncanonical residue side chain. **DTTA-C3-yne**, which has a shorter linker, showed a larger $\Delta\chi$ tensor than did **DTTA-C4-yne**. **DTTA-C3-yne** was used in the structure determination of a transient protein complex.²⁵⁹ Chen et al. reported phenyl-sulfonated pyridine modified DTTA probes (Figure 7, **71–72**), which can react with cysteine with formation of a thioether bond shown to be stable in a cell lysate.²⁶⁰ **4PS-6M-PyDTTA** (Figure 7, **72**) has an extra methyl group in position six of the pyridine ring. This small difference proved to alter the speed of probe ligation, lanthanoid ion binding affinity, and $\Delta\chi$ tensor. **4PS-6M-PyDTTA** showed a higher reactivity toward ubiquitin S57C and the *Staphylococcus aureus* sortase A (SrtA) mutant D82C.²⁶⁰

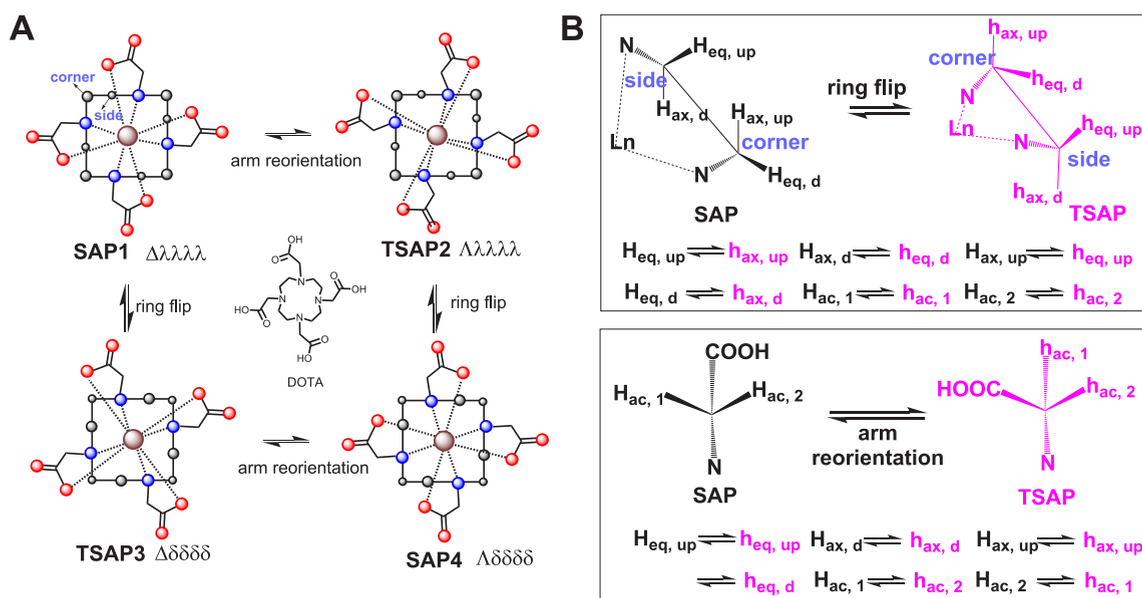


Figure 8. (A) Schematic representation of four different stereoisomers populated by DOTA lanthanoid complexes and their conformational exchange. The cyclen ring is drawn as a square with solid lines; nitrogen, oxygen, and carbon atoms are shown as blue, red, and black spheres, respectively; the metal ion is represented by a brown sphere. (B) Newman projections showing the effect of the ring flip on the positions of the hydrogen atoms of two neighboring carbon atoms. Adapted from Q. Miao (2019) Design, synthesis and application of paramagnetic NMR probes for protein structure studies. PhD Thesis, Leiden University.

Following attachment, **6M-PyDTTA** exhibited a lower metal binding affinity but larger $\Delta\chi$ tensor than PyDTTA. Interestingly, the two probes loaded with Tm(III) gave PCSs of opposite sign, indicating different orientations of the $\Delta\chi$ tensor.²⁶⁰

3.3.2.3. PyMTA-Based Probes. In contrast to the linear structure of EDTA, 2,2',2'',2'''-(pyridine-2,6-diylbis(methylene))bis(azanetriyl)tetracetic acid (PyMTA) has a pyridine core with diazandiylacetic acid pendants in positions 2 and 6. Its Ln(III) complexes have been used for luminescence experiments and as contrast agents in MRI.^{261,262} **4MMPyMTA** (Figure 7, 73) was designed for chemical stability and successfully used as a paramagnetic lanthanoid probe to study the structures of minor protein species.²⁶³ **4V-PyMTA** (Figure 7, 74)²⁶⁴ was reported as a paramagnetic Ln(III) probe with a vinyl group in position 4, which forms a thioether bond with cysteine. After successful attachment of **4V-PyMTA** to ubiquitin, small PCSs were observed.²⁶⁴ Following substitution of position 3 of the pyridine ring with bromide to obtain probe **75**, the reactivity was improved, but not the paramagnetic effects.²⁵ Further work introduced phenylsulfonated in pyridine position 4 as the protein conjugation group (Figure 7, 76).²⁶⁵ Compared with probes 73 and 74, probe **76** yielded larger PCSs due to the short and rigid thioether attachment, but with a low reaction rate, even under fairly harsh conditions.²⁶⁵ Subsequently, probe **77** (Figure 7)²⁶⁶ was constructed, which features both good reactivity and selectivity for solvent exposed cysteine residues.

3.3.2.4. General Synthetic Strategies of Aminopoly-(Carboxylic Acid)-Based Probes. There are two main strategies for the synthesis of aminopoly(carboxylic acid) based probes. The first involves either the direct modification of the carboxylic acid with attachment groups by a coupling reaction, such as applied for probes **60** (derived from EDTA) and **66** (derived from DTPA), or functionalization of the central amine with an attachment group as in, for example, probes **69** and **70** (derived

from protected DTTA). The second strategy starts from the core structure of the probe, introduces the attachment group, and subsequently modifies the intermediate with a polyamine to introduce the carboxylic acid pendants. For example, the synthesis of probes **71** and **72** starts from a mono- or dimethylpyridine derivative, to which the attachment group (benzenesulfonyl) is added after several modification steps. Thus, after N-oxidation, nitration, rearrangement to their trifluoroacetate esters, and in situ mild hydrolysis of α -picoline or 2,6-lutidine, a nucleophilic aromatic substitution provided the sulfones. By alkylation with a halogenated pyridine derivative (Scheme 7, structure colored in red), the polyamine intermediate (Scheme 7, structure colored in magenta) is generated after deprotection and finally the poly(carboxylic acid)s are introduced to give the probe (Scheme 7, structure colored in blue).²⁶⁰

3.3.3. Cyclen-Based Tags. Cyclen(1,4,7,10-tetraazacyclododecane) is a cyclic ring molecule of tetraethyleneamine, which can be modified to obtain various metal binding complexes, which are particularly suited for lanthanoid ions. DOTA (1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid) is one of the most frequently used cyclen derivatives. It was first synthesized in 1976.²⁶⁷ DOTA complexes are thermally stable and bind Ln(III) ions tightly.²⁶⁸ It is important to note that Ln(III)-DOTA complexes undergo conformational exchange involving cyclen ring flips combined with reorientation of the tetracarboxylic acid pendants (Figure 8). The pendant arms can coordinate clockwise (Λ) or anticlockwise (Δ) and the cyclen ring has two conformations ($\lambda\lambda\lambda\lambda$ and $\delta\delta\delta\delta$). Consequently, Ln(III)-DOTA forms two pairs of enantiomeric species (square antiprism, SAP, and twisted square antiprism, TSAP) in solution, which tend to produce different magnetic effects in the case of anisotropic paramagnetism and averaging between them due to conformational exchange.^{269–271} Therefore, the designs of the cyclen probes described below aim to lock the cyclen ring in a single conformation to obtain uniform

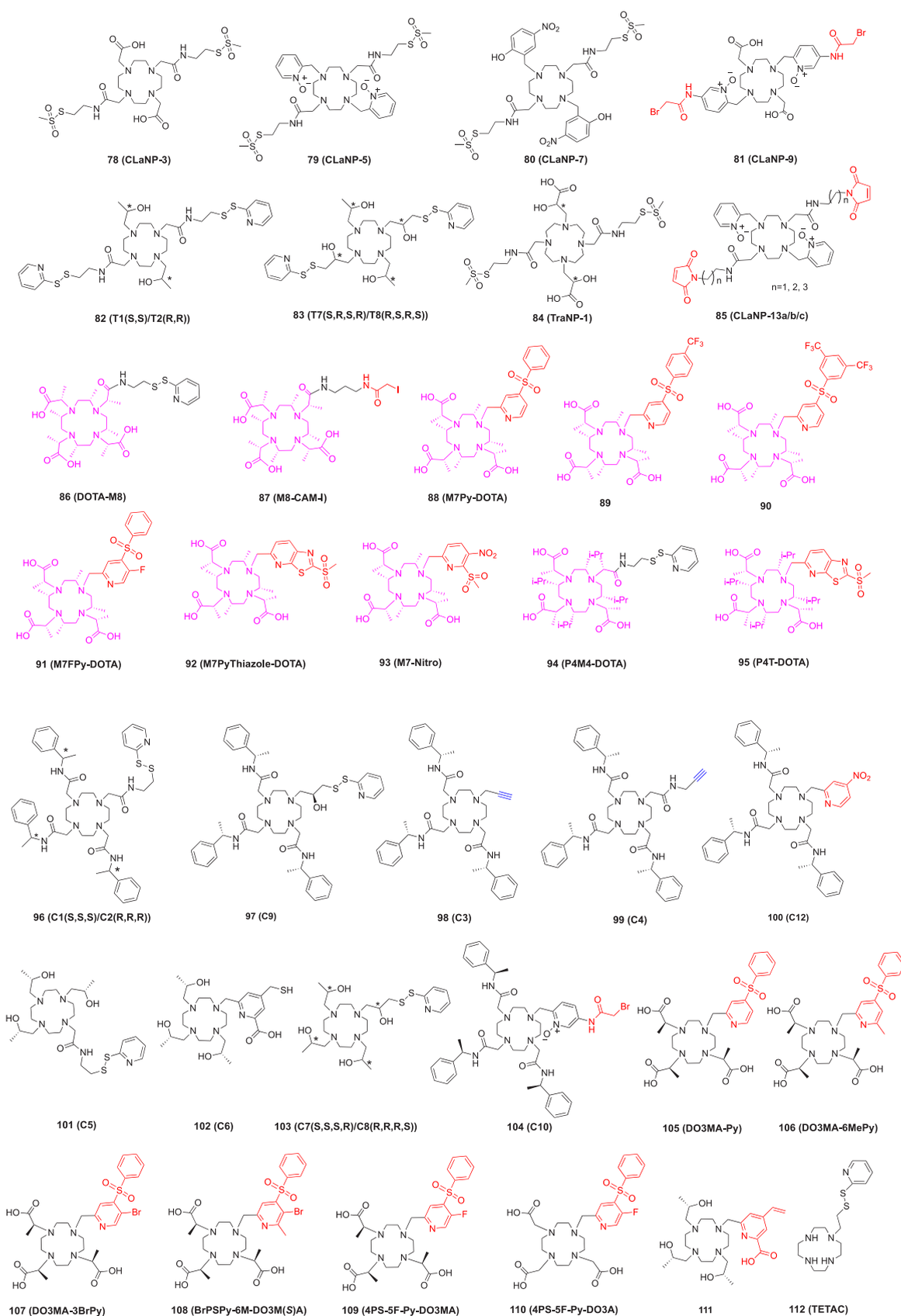


Figure 9. Chemical structures of cyclen-based paramagnetic probes. The functional groups for attachment to cysteine via thioether formation are highlighted in red. The functional groups for attachment to an azide group in nCAA via “click” reaction are highlighted in blue. The cyclen rings with substituents are shown in magenta. 78,²⁷² 79,^{127,273} 80,²⁷⁶ 81,²⁷⁷ 82,²⁷⁸ 83,²⁷⁹ and 84²⁸⁰ are double-armed probes. Asterisks identify chiral carbon atoms.

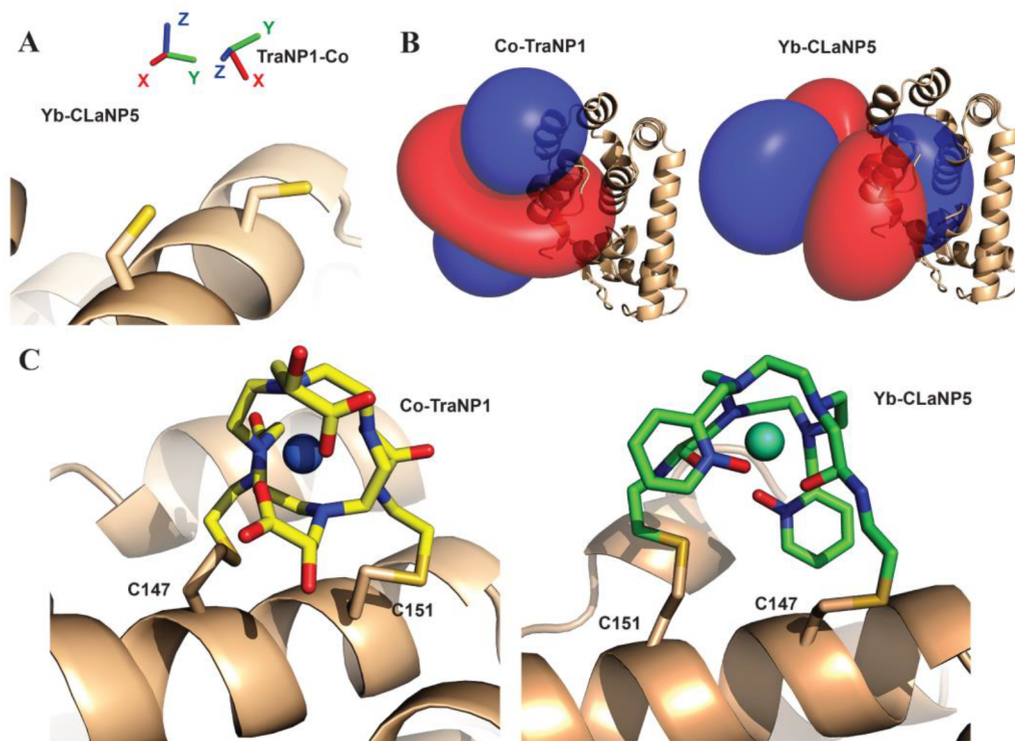


Figure 10. Comparison of Co(II)-TraNP-1-SS and Yb(III)-CLaNP-5 attached to T4Lys K147C/T151C. (A) Metal positions and tensor orientations. (B) PCS iso-surfaces (± 0.4 ppm for Yb(III)-CLaNP-5, ± 0.2 ppm for Co(II)-TraNP-1-SS). (C) Models of the protein–probe structures. The protein is shown in ribbon representation; the probes and cysteine residues are shown as sticks and the metal ions as spheres. Reproduced without changes from ref 280. Copyright 2019 the authors, under CCA license (<https://creativecommons.org/licenses/by/4.0/>).

paramagnetic effects, by incorporating substituents on the cyclen ring and/or the pendant arms with the appropriate chirality.

3.3.3.1. Double-Anchored Attachment. The first reported cyclen based protein paramagnetic probe was CLaNP-3 (Figure 9, 78)²⁷² in complex with paramagnetic Ln(III) ions. By functionalizing two opposing tetracarboxylic acid arms of DOTA with MTS groups, CLaNP-3 can be anchored to a protein via two disulfide-bridges for maximum immobilization of the metal ion relative to the protein. For pseudoazurin (Paz) with mutations E51C and E54C tagged with Yb(III)-CLaNP-3, PCSs were detected for nuclear spins up to 35 Å from the paramagnetic center. Some residues, however, displayed peak doubling, suggesting that the complex populated more than a single conformation with different $\Delta\chi$ tensors.²⁷²

Subsequently, CLaNP-5 (Figure 9, 79)^{127,273} was produced and became one of the most used lanthanoid probes. In CLaNP-5, the two acetate pendant arms of CLaNP-3 were replaced by pyridyl-*N*-oxides, which disfavors the population of the TSAP conformation of its lanthanoid complexes. For Yb(III)-CLaNP-5 linked to Paz E51C/E54C, a single set of large PCSs was observed. In contrast, CLaNP-5 attached via a single arm yielded only small PCSs due to averaging caused by the flexibility of the attachment.²⁷³ Further studies demonstrated that magnetic properties of CLaNP-5 are insensitive to the chemical environment of the attachment site, allowing the prediction of the $\Delta\chi$ tensor parameters and Ln position for a known protein structure with reasonable accuracy.¹²⁷ CLaNP-5, thus, is highly suitable for structural studies of proteins and protein complexes. It turned out, however, that the detection of little populated protein states by relaxation dispersion (RD) experiments is

compromised by the presence of minor conformations of the complex that can lead to probe-induced RD effects.^{274,275} Other disadvantages of CLaNP-5 are its charge (+3), making it less favorable for some proteins and protein complexes, and the disulfide bonds formed with the protein are liable to reduction, which can lead to loss of the probe over a period of several weeks.

To decrease the charge, CLaNP-7 (Figure 9, 80)²⁷⁶ contains two phenolic groups, lowering the net charge to +1 by deprotonation of the phenol hydroxyls. CLaNP-7 is yellow in color, which is convenient during the tag labeling reaction and protein purification. Similar to CLaNP-5, it induced a single set of PCSs in Paz E51C/E54C, with a $\Delta\chi$ tensor that was only slightly smaller than that of CLaNP-5.²⁷⁶ Unexpectedly, two sets of PCSs were observed specifically for Yb(III)-CLaNP-7 linked cytochrome *c* and this peak doubling was pH-dependent. By mutation of His39, which is adjacent to the probe attachment site, to alanine, it was demonstrated that the peak doubling was caused by an interaction with this residue, suggesting that the hydration water molecule, which coordinates the Ln(III) ion in its ninth coordination site, forms a hydrogen bond with the imidazole ring of His39 and in this way breaks the C_2 symmetry of the probe. The effect was pH dependent due to protonation and deprotonation of the imidazole ring. Since disulfide bonds can easily be reduced, CLaNP-9 (Figure 9, 81)²⁷⁷ was designed as a variant of CLaNP-5, featuring two bromoacetanilide groups to react with cysteine residues by formation of chemically stable thioether bonds. Using T4 lysozyme (T4Lys) N55C/V57C and Paz E51C/E54C as model proteins, Yb(III) loaded CLaNP-9 yielded single sets of PCSs under reducing conditions, indicating the high stability of this probe. Because of the rapid hydrolysis of

the bromoacetanilide groups in aqueous solutions, however, quantitative tagging proved difficult to achieve.

The tags **T1** and **T2** (Figure 9, 82)²⁷⁸ similarly are double-anchored cyclen based probes using pyridylthio groups as attachment groups. **T1** and **T2** are enantiomers, as their two free coordination arms are chiral isopropanol moieties. Large $\Delta\chi$ tensors were found for these lanthanoid complexes linked to *Staphylococcus aureus* 6-hydroxymethyl-7,8-dihydropterin pyrophosphokinase (HPPK) K76C/C80, and exceptionally narrow distance distributions were obtained by DEER measurements using the tags loaded with Gd(III).²⁷⁸ The paramagnetic effects induced by other paramagnetic Ln(III) ions, however, are sensitive to the pH, with large PCSs and RDCs detected at high pH. Another pair of enantiomeric probes, **T7** and **T8** (Figure 9, 83),²⁷⁹ were designed with two shorter chiral attachment arms. Although large $\Delta\chi$ tensors were determined for a wide range of pH values for both tags, the χ tensor orientation changed between high pH (8) and low pH (6.5). It was noted that **T7** produced incomplete ligation yields for one of the test proteins (HPPK K76C/C80).²⁷⁹

The tags described above are all lanthanoid based cyclen probes. Recently, several double-arm 3d block ion probes (transition metal ion NMR probes, TraNPs) were reported.²⁸⁰ Among them, **TraNP-1** (Figure 9, 84)²⁸⁰ can tightly bind Co(II) and Mn(II) ions. Its Co(II) complex generates a single set of PCSs and yields a medium sized tensor, which is suitable for application in small proteins or localized studies in larger proteins or protein complexes. The authors compared the tensor orientation between Co(II)-**TraNP-1** and Yb(III)-**CLaNP-5** ligated to T4Lys K147C/T151C by $\Delta\chi$ tensor fits using experimentally measured PCSs. The tags delivered significantly different tensor orientations as evidenced by the PCS iso-surfaces shown in Figure 10.²⁸⁰

Recently, the double-anchored probe **CLaNP-13** (Figure 9, 85) was designed for DEER studies in living cells.²⁸¹ By virtue of two maleimide groups, **CLaNP-13** forms thioether bonds with cysteine residues, which in a cellular environment are more stable than disulfide bonds. As Michael additions of thiols to maleimides generate enantiomers, NMR measurements showed peak doubling for the Yb(III) complex attached to T4Lys double-cysteine mutants. DEER measurements, however, yielded a narrow distance distribution for **CLaNP-13** attached to the quadruple mutant T4Lys N55C/V57C/K147C/T151C, and the probe was also shown to be suitable for in-cell DEER measurements.

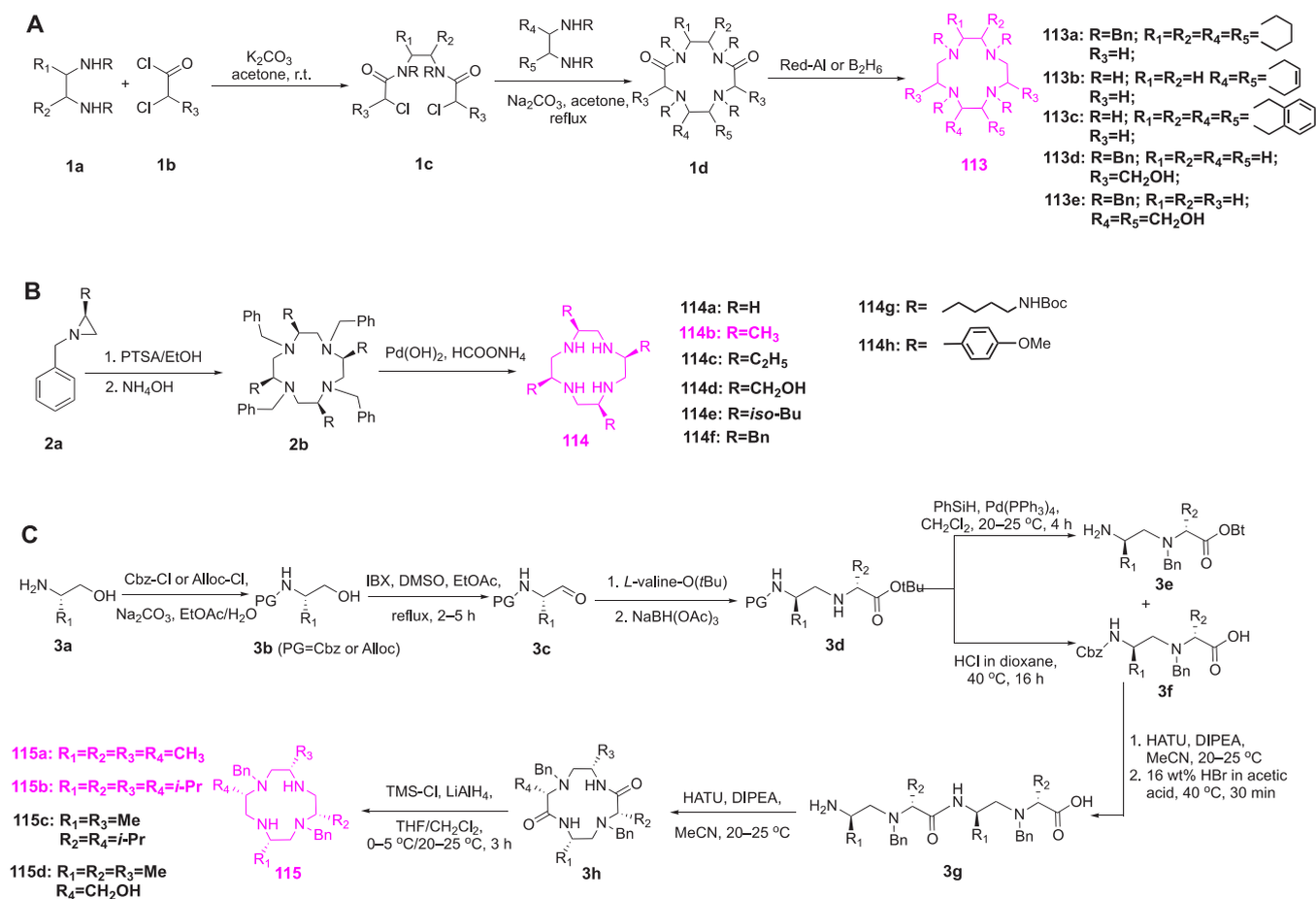
3.3.3.2. Single Site Attachment. **DOTA-M8** (Figure 9, 86)²⁸² is a single-arm lanthanoid probe, which produces $\Delta\chi$ tensors of remarkable size. Its structure is based on M4DOTMA.²⁸³ Because of the eight chiral methyl substitutions on the cyclen ring and acetate pendant arms, M4DOTMA loaded with Yb(III) exhibits a single conformation, which is rigidified by steric crowding.²⁸³ A set of large PCSs was reported for **Dy(III)-DOTA-M8** labeled ubiquitin S57C. **Dy(III)** is tightly bound to **DOTA-M8**, withstanding harsh chemical and physical conditions.²⁸² About 15–20% of a minor species appeared in the NMR spectra, however, and this ratio increased to 50% by heating the tagged protein to 323 K. Subsequent studies revealed slow conformational exchange between two coordination geometries of **Ln(III)-DOTA-M8**, which produce different $\Delta\chi$ tensors, with the exchange rate depending on the radius of the lanthanoid ion.^{271,284}

As **DOTA-M8** is ligated to the protein via a disulfide bond, the ligation is not stable under reducing conditions. This prompted

efforts to link **DOTA-M8** to proteins by different attachment chemistries. By replacing the pyridylthio linker arm in **DOTA-M8** by a propylidoacetamide ($-C_3H_6NHCOCH_2I$) group, a chemically more stable thioether bond is formed in the reaction with cysteine. This new tag was referred to as **M8-CAM-I** (Figure 9, 87) and its usefulness for in-cell NMR measurements was demonstrated, but the longer linker resulted in a smaller $\Delta\chi$ tensor than for **DOTA-M8**.²⁸⁵ By using 4-(phenylsulfonyl)pyridine as the attachment group, another **DOTA-M8** derivative, referred to as **M7Py-DOTA** (Figure 9, 88), forms a more rigid and shorter linkage to the protein, which is also suitable for in-cell NMR.²⁸⁶ As the reactivity of the 4-(phenylsulfonyl)pyridine group is relatively low, the CF_3 -substituted phenyl analogues (Figure 9, 89 and 90) were designed for enhanced reactivity, but this approach did not increase the rate of protein tagging because the rate-limiting step appears to be the formation of a Meisenheimer-type complex rather than the dissociation of the pyridine-sulfone moiety.²⁸⁷ Consequently, the probes **M7FPy-DOTA** (Figure 9, 91)²⁸⁷ and **M7PyThiazole-DOTA** (Figure 9, 92)²⁸⁷ were produced, both of which proved to react with cysteine with high selectivity and efficiency. The authors found that **M7FPy-DOTA** can react also with tris(2-carboxyethyl)phosphine (TCEP) to release benzenesulfonic acid, which can catalyze protein dimerization by intermolecular disulfide bridge formation between cysteine residues. **M7PyThiazole-DOTA** performs better, as it is more reactive toward cysteine than TCEP and generates an even larger $\Delta\chi$ tensor than **DOTA-M8**.²⁸⁷ Recent work introduced a 3-nitro-substituted 2-(methylsulfonyl)pyridine moiety as attachment group to design a new **DOTA-M8** based probe named **M7-Nitro** (Figure 9, 93).²⁸⁸ This probe possesses high protein ligation rates (more than 95% ligation in 30 minutes at room temperature) and a very large $\Delta\chi$ tensor ($\Delta\chi_{ax} = 94.3 \times 10^{-32} \text{ m}^3$) was measured with **Dy(III)-M7-Nitro** labeled ubiquitin S57C. The authors attributed the improvements in ligation to the methylsulfone leaving group in the *ortho* position and the presence of the nitro substituent in the *meta* position.

To further improve the rigidity of the cyclen-based probes, **DOTA** derivatives with more crowded substituents, **P4M4-DOTA** (Figure 9, 94)¹²⁶ and **P4T-DOTA** (Figure 9, 95)²⁸⁹ were synthesized. Only a single conformation, $\Lambda(\delta\delta\delta\delta)$, was detected in ROESY spectra of the **Lu(III)-P4M4-DOTA** complex.¹²⁶ Linking lanthanoid ion loaded **P4M4-DOTA** to ubiquitin S75C and human carbonic anhydrase II (hCA II) S166C delivered single sets of PCSs, but some of the *Q* factors of the $\Delta\chi$ tensor fits were relatively large owing to the length and flexibility of the attachment arm. As expected, **P4T-DOTA** performs better than **P4M4-DOTA** because of a shorter and more rigid linker. **P4T-DOTA** also forms a stable thioether bond with cysteine.

C1 (Figure 9, 96) is another early example of a **DOTA** based probe for protein studies by paramagnetic NMR, which is relatively easy to synthesize.²⁹⁰ In this tag, the metal ion is coordinated by amides instead of carboxyl groups. Loading of **C1** with Ln(III) ions requires harsh conditions and the ions are tightly bound thereafter. Three chiral bulky pendant arms ensure that the **C1** tag populates a single conformation as indicated by ¹H NMR spectra of Yb(III) loaded **C1** and single sets of PCSs were observed with all proteins tested. **C2** (Figure 9, 96) is the enantiomer of **C1** and generates a slightly different tensor orientation.²⁹¹ On the basis of the structure of probe **C1**, the protein EPR probe **C9** (Figure 9, 97)²⁹² was designed with a chiral tagging arm. Variants of **C1** with alkyne attachment

Scheme 8. Three Strategies for Cyclen Ring Modulation^a

^aThe magenta colored structures show the final cyclized ring with different substituents. The compounds in magenta are applied for paramagnetic probe design.

groups, C3 and C4 (Figure 9, 98, 99),²⁹³ were developed for ligation to the ncAA AzF (Figure 4, 9a) by “click” chemistry.²⁹³ The PCSs obtained with probe C4 are smaller than those of probe C3 due to the longer linker of the former. The Cu(I) catalyzed ligation reaction was shown to be selective also in the presence of cysteine residues in the target protein and thus suitable for proteins containing cysteine residues of structural and functional importance. Subsequent experience showed, however, that many proteins are prone to precipitation in the presence of Cu(I) catalysts.¹⁸² For this reason, the recently published C12 probe (Figure 9, 100)²⁹⁴ reacts with cysteine by formation of a thioether bond to its aromatic ring linkage group. Complete tagging yields of cysteine were obtained at room temperature overnight. In contrast, the tag reacts with selenocysteine in minutes, providing a route to selective tagging of proteins in the presence of native cysteine residues.²⁹⁴

Ln(III) complexes of C1 and its variants are relatively large, and although experimental evidence is scarce,²⁹⁵ their hydrophobic aromatic rings may result in unexpected specific interactions with the protein surface. Therefore, a number of alternative cyclen probes were produced with hydrophilic pendants, where the single conformation of the cyclen ring is ensured by the chirality of the pendant arms. The probe C5 (Figure 9, 101)¹²⁹ contains three chiral (*S*)-2-hydroxypropyl groups as pendant arms. Variants of C5 with different attachment linkers were also explored, including C6 (Figure 9,

102),¹²⁹ which contains a rigid picolinic acid linker, C7 (Figure 9, 103),¹²⁹ which contains a chiral alcohol in the attachment arm instead of an amide, and C8 (Figure 9, 103),¹²⁹ which is the enantiomer of C7. Like C1 and C2, these tags can generate large $\Delta\chi$ tensors. A comparison of three proteins and different attachment sites showed, however, that the $\Delta\chi$ tensor produced by C7 is highly sensitive to the protein environment, which is unusual for lanthanoid based cyclen probes.¹²⁹ Probe C10 (Figure 9, 104)²⁹⁶ was designed for DNA studies (discussed in section 3.4). Ligated to ubiquitin A28C, it produced $\Delta\chi$ tensors of medium size, as expected for the relatively long and flexible attachment arm. Recently, DO3MA-Py (Figure 9, 105),²⁹⁷ DO3MA-6MePy (Figure 9, 106),²⁹⁷ DO3MA-3BrPy (Figure 9, 107),²⁹⁸ and BrPSPy-6M-DO3M(S)A (Figure 9, 108)²⁹⁹ were reported, which contain methyl substituted carboxylic acid pendants and phenylsulfonated pyridine with different substituents on the pyridine ring as functional group for attachment. These lanthanoid ion complexes carry zero net charge and form thioether bonds with cysteine residues. Large PCSs were observed with these probes. While the ligation rates of DO3MA-Py and DO3MA-6MePy are slow even at high pH and temperature, the bromide substituent in DO3MA-3BrPy decreased the reaction time to 6 h at ambient temperature. Sizeable $\Delta\chi$ tensors were obtained with the Yb(III) complex of DO3MA-3BrPy attached to ubiquitin D39C and ubiquitin E64C, and DO3MA-3BrPy loaded with Gd(III) was also

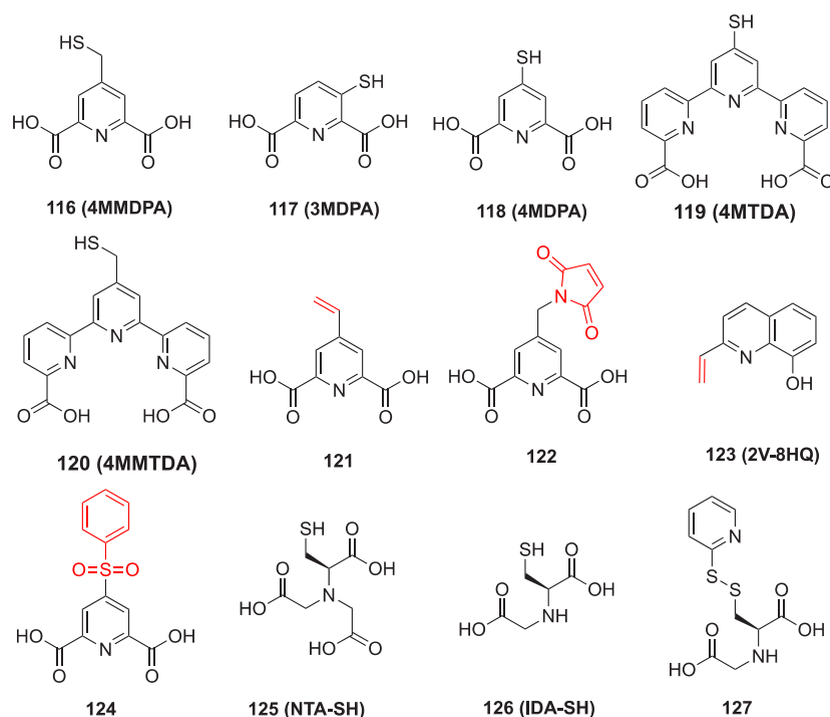


Figure 11. Chemical structures of small molecule probes. Functional groups designed for attachment via thioether formation are highlighted in red. 116,³²² 117,³²³ 118,³²⁴ 121,³²⁶ 122,³²⁶ and 124²⁶⁵ are DPA-based probes. 119 and 120 are TDA based probes. 123¹⁷⁶ is based on 8-hydroxyquinoline. 125³²⁷ is an NTA based probe. 126³²⁸ and 127³²⁹ are IDA based probes.

successfully used for in-cell distance measurements by EPR, although the distance distribution obtained was relatively broad.²⁹⁸ Its companion **BrPSPy-6M-DO3M(S)A** has an extra methyl group on the pyridine and the authors reported that this design improved the rigidity of the tag.³⁰⁰ By replacing bromine in **DO3MA-3BrPy** for a fluorine atom, Su and co-workers synthesized the probes **4PS-5F-Py-DO3MA** (Figure 9, 109) and **4PS-5F-Py-DO3A** (Figure 9, 110) to analyze the conformational exchanges of DOTA-Ln-like probes using 1D and 2D EXSY NMR spectroscopy.³⁰¹ They demonstrated that even the presence of chiral centers in three of the pendants of a DO3A-type cyclen tag does not prevent the population of a minor conformational species, which persists following ligation of the lanthanoid complexes to ubiquitin G47C.³⁰¹ Notably, a second conformational species has also been reported for **DOTA-M8**, although this probe contains four additional stereocenters in the cyclen ring.²⁸⁷ The relative conformational stabilities of SAP and TSAP conformations of DOTA-type cyclen-Lu(III) complexes with different chiralities of methylated cyclen and carboxylate pendants has recently been rationalized computationally.³⁰²

Another recent approach used a photocatalyzed reaction to conjugate a vinyl probe to proteins.³⁰³ Using 2,2-dimethoxy-2-phenylacetophenone (DPAP) as a photoactivated radical initiator, the ligation rate of the vinylpicolinic acid modified probe 111 (Figure 9) with GB1 T53C was greatly accelerated, but the $\Delta\chi$ tensors obtained were small.³⁰³

3.3.3.3. General Synthetic Approaches of Cyclen Based Probes. DOTA and its analogues have been used as metal ion ligands for decades and the thermodynamic properties, synthesis, and applications of their metal complexes have been investigated in detail.^{268,304–308} Commensurate with the wide range of applications for cyclen and DOTA compounds in many different fields, numerous synthetic routes have been reported. It

should be noted that DOTA based probes with chiral substituents to ensure conformational rigidity, require the execution of a multistep synthetic scheme. In addition, the protective group strategy has to be chosen carefully to allow the introduction of one or two attachment arms. Among the different synthetic methods reported for large-scale cyclen preparation,^{307,309–314} three main strategies stand out for modification of the cyclen ring. The first is based on one of the synthetic procedures of cyclen (Scheme 8A). This approach was explored by Jacques et al., who used substituted ethylenediamine derivatives and chloroacetyl chloride derivatives as starting materials to form a linear diamide intermediate (**1c** in Scheme 8A), which was further reacted with an ethylenediamine derivative to yield the cyclic product **1d**. Finally, the amide groups of the ring were reduced to give the core macrocyclic structure **113** (Scheme 8A).^{315,316} On the basis of this method, symmetric and asymmetric functionalized cyclen derivatives were produced, such as cyclohexyl, cyclohexene and tetraline substituted cyclens **113a–c**,³⁰⁴ and the diol substituted compounds **113d–e**.^{315,316} Later on, the same group developed a new method to synthesize cyclen derivatives with C4 symmetry, where, instead of using a stepwise cyclization, a more straightforward synthetic route was followed by oligomerization of *N*-substituted aziridine (**2a** in Scheme 8B) in the presence of *p*-toluenesulfonic acid (PTSA, Scheme 8B).^{283,317} This method directly gives the *N*-protected cyclen derivatives (**2b** in Scheme 8B), albeit in low yield, and, after deprotection, the desired macrocycle **114**. The cyclen based probes **DOTA-M8**, **M7Py-DOTA**, **M7FPy-DOTA**, **P4T-DOTA** as well as **M7PyThiazole-DOTA** (Figure 9) were synthesized using this approach and several other groups subsequently used this approach for cyclen based ligand synthesis.^{313,314} Anelli et al. employed both approaches for the synthesis of more hydrophilic macrocyclic compounds.³¹⁸

Recently, Häussinger and co-workers reported a new method for synthesizing modified cyclens (Scheme 8C),^{126,319} which starts from chiral aminoaldehydes (3c in Scheme 8C) derived from their respective amino acid precursors. Via reductive amination, two suitably protected linear alkylated amino acid derivatives (3e and 3f) are produced, which are readily coupled to each other and cyclized by standard solution peptide coupling under high dilution. Compared to the other two strategies, this newly reported route provides a high yield in the cyclization step and the overall yield of macrocycles 115 has been reported around 22%.³¹⁹

3.3.4. Small Molecule Tags. Small molecules like dipicolinic acid (DPA), iminodiacetic acid (IDA), nitrilotriacetic acid (NTA), and [2,2':6',2'']terpyridine-6,6''-dicarboxylic acid (TDA) have also been explored for the design of paramagnetic NMR probes. Their attraction derives from their small size, which minimizes the potential impact on the protein structure, and their relative ease of synthesis. In complexes with Ln(III) ions, these small probes leave several coordination sites of the Ln(III) ion free for coordination by solvent molecules or additional protein side chains. With these tags, the metal complexes are formed by titration with metal ions after probe attachment, which opens the possibility of inaccurate metal to protein ratios and binding of metals to other, undesired sites.

3.3.4.1. DPA, TDA, and HQA Tags. DPA coordinates metal ions in nonchiral geometry, thus reducing the potential for different enantiomeric forms of metal coordination and peak doubling in NMR spectra. Despite presenting only a tridentate ligand, the affinity of DPA for lanthanoid ions is in the low nanomolar range.^{320,321} 4MMDPA (Figure 11, 116)³²² was the first DPA tag developed. It was attached to a cysteine residue following activation with DTNB, which results in a disulfide linkage, and shown to generate PCSs following attachment to Cys68 of the ArgN protein, in which the coordination of the paramagnetic lanthanoid ion appears to be supported by coordination to a glutamate side chain.³²² Despite the limited coordination of the lanthanoid ion, the dissociation rate of the metal ion was slow at 25 °C (0.1 s⁻¹), but the $\Delta\chi$ tensor was smaller than that found with cyclen based tags. Loaded with Gd(III) ions, the tag was also demonstrated to be suitable for Gd–Gd distance measurements by EPR spectroscopy in different proteins.¹²² To shorten the linker length, the probes 3MDPA (Figure 11, 117)³²³ and 4MDPA (Figure 11, 118)³²⁴ were produced, which feature the sulfur atom right on the pyridine ring. 3MDPA, which has a thiol group in position 3, was ligated to ArgN like 4MMDPA via a disulfide bond following activation of its cysteine residue with DTNB. In complex with paramagnetic Ln(III) ions, $\Delta\chi$ tensor orientations were obtained different from those with 4MMDPA and metal coordination still appeared to be assisted by Glu21, but the magnitudes of the $\Delta\chi$ tensors were small. Compared to the 3MDPA tag, the disulfide dimer of 4MDPA is a more convenient tag for ligation, as it reacts spontaneously and quantitatively with cysteines by disulfide exchange. While it displayed relatively small $\Delta\chi$ tensors with ArgN, larger $\Delta\chi$ tensors were obtained with the intracellular domain of the p75 neurotrophin receptor (p75 ICD). The tensor orientations differed substantially from those obtained with the corresponding 4MMDPA complexes for ArgN but not for p75 ICD, suggesting important contributions by additional coordinating amino acid side chains. The DPA tags also display good affinity for Co(II), as demonstrated first for ArgN with 3MDPA tag.³²³

Using DPA as starting materials, TDA based probes, 4MTDA (probe 119, Figure 11)³²⁵ and 4MMTDA (probe 120, Figure 11)³²⁵ were synthesized. Large PCS were detected after their Ln ion loaded complexes were ligated to ubiquitin T22C or A28C. They also exhibit fluorescent properties. For both mutants, ubiquitin side chains are involved in metal coordination. The authors also showed that 4MTDA exchanges lanthanide ions more slowly than DPA, as 2D ¹H–¹H EXSY spectra showed no observable chemical exchange. The ligation yields were reported to be relatively low.

Probe 121 (Figure 11)³²⁶ contains a vinyl group to link the probe to the protein via Michael addition to form a thioether linker. The vinyl group avoids the problem of maleimide functionalized tags (Figure 11, 122), which create a new chiral center upon reaction with a thiol group.³²⁶ A vinyl group was also used to ligate the 8-hydroxyquinoline based probe 2V-8HQ (Figure 11, 123)¹⁷⁶ to a cysteine. While the tag is not suitable for binding lanthanoid ions, it binds divalent transition metal ions such as Mn(II), Co(II), Ni(II), Cu(II), and Zn(II) with micromolar affinities, and a large $\Delta\chi$ tensor was observed for the Co(II) complex of 2V-8HQ bound to ubiquitin A28C/E24H, where the binding of the metal ion was assisted by the histidine residue.¹⁷⁶ A DPA derivative with a phenylsulfonated pyridine functional group (Figure 11, 124)²⁶⁵ was shown to generate a stable, rigid and short tether with a thioether bond to a cysteine, but $\Delta\chi$ tensor parameters were not reported.

3.3.4.2. NTA and IDA Tags. Similar to DPA and 8HQ tags, NTA-based probes immobilize metal ions on target proteins depending on the assistance from additional coordinating groups provided by other amino acid residues. The first NTA-based probe suitable for binding lanthanoid ions, NTA-SH (Figure 11, 125),³²⁷ was obtained by modifying cysteine with two acetate groups. Single sets of PCSs were reported upon complexation with paramagnetic lanthanoid ions following attachment of the probe to either one or two cysteine residues of the target protein. For a single NTA-SH tag, much larger $\Delta\chi$ tensors were observed with ArgN than ubiquitin A28C, suggesting that the protein environment plays a critical role in immobilizing the Ln(III) ion. This difficult-to-predict variability can be reduced by simultaneously attaching NTA-SH tags at positions of *i* and *i* + 4 of an α -helix to create a metal binding site with coordination by both NTA moieties. This approach can also deliver very large $\Delta\chi$ tensors.³²⁷ As the metal ion affinity is not as high as for DOTA ligands, the metal ion can be exchanged by washing with EDTA and subsequently introducing another metal ion.

An even smaller tag is based on IDA, which only contains two carboxylates. The IDA-SH tag (Figure 11, 126)³²⁸ and its activated form (Figure 11, 127)³²⁹ were reported to yield a single set of PCSs after chelating lanthanoid ions. This tag can yield large $\Delta\chi$ tensors, if the Ln(III) ion can be additionally coordinated by a protein side chain as observed for ubiquitin A28C, where Asp32 is located in the same α -helix as residue 28.³²⁸ By ligating two IDA-SH tags to cysteine residues in positions *i* and *i* + 4 of an α -helix, a Co(II) ion can be coordinated in an octahedral complex and produces sizable PCSs (Figure 11, 127).³²⁹ In ubiquitin E24C/A28C, however, this tag arrangement performed best with substoichiometric quantities of Co(II) and Zn(II) ions, whereas lanthanoid ions produced multiple species. Subsequent study of this tag and metal complexation mode in the same protein and in T4Lys K147C/T151C revealed additional species also with Co(II) ions and difficulties to achieve complete saturation of the metal

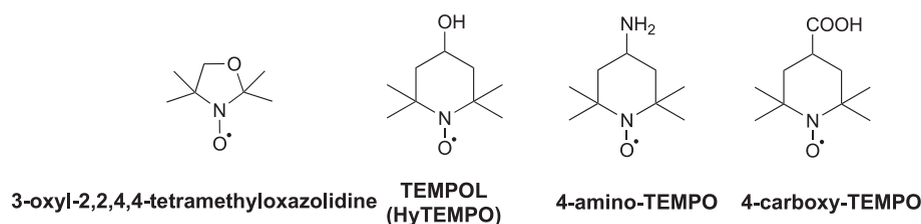


Figure 12. Chemical structures of nitroxide based cosolute probes.

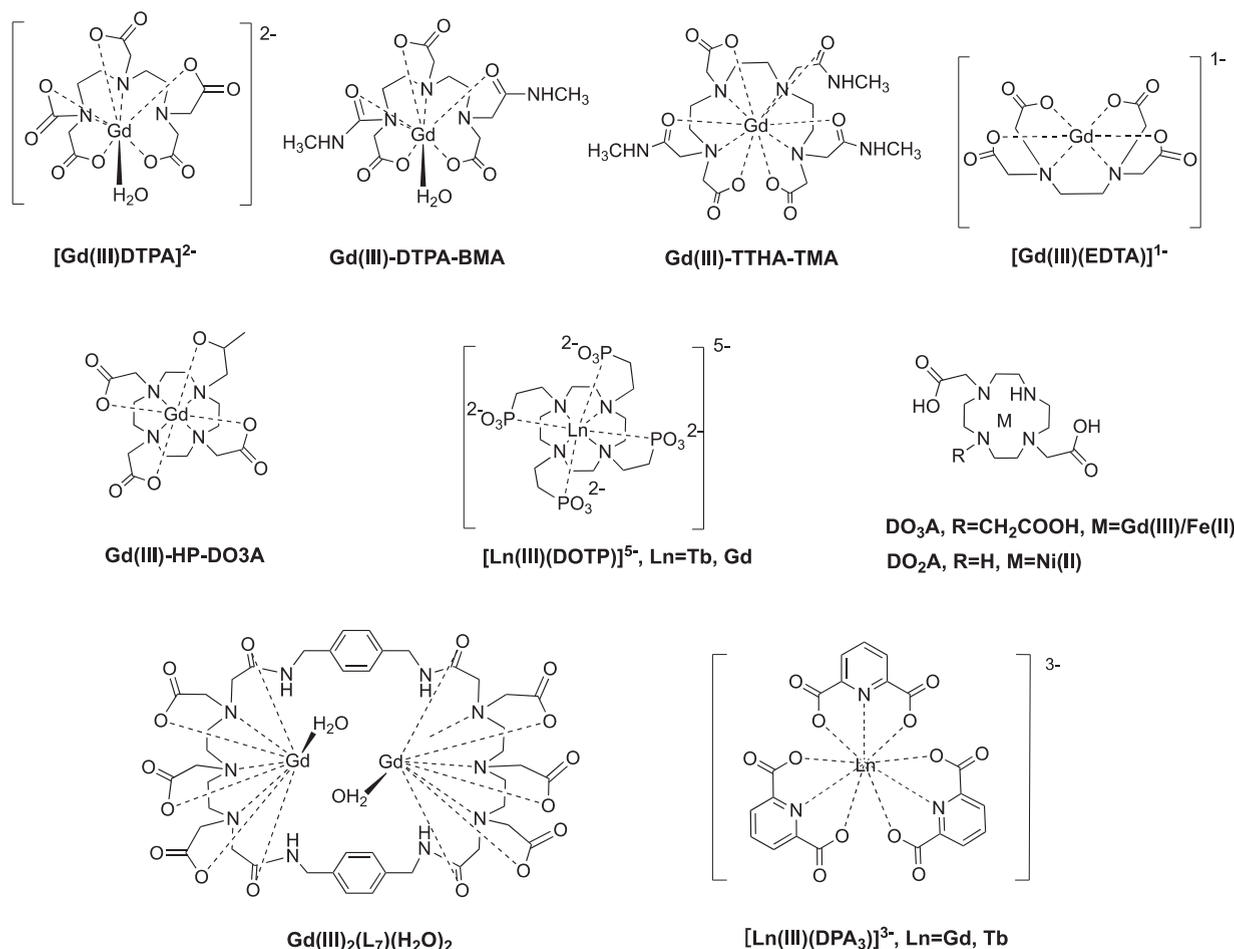


Figure 13. Chemical structures of metal ion based cosolute probes.

binding site.²⁸⁰ A systematic exploration of the best binding partner for an **IDA-SH** or **NTA-SH** tag attached to a cysteine residue in position i of an α helix indicated that the **IDA-SH** tag forms the best Ln(III) binding sites together with an aspartate residue in position $i + 4$, while the sterically more demanding **NTA-SH** tag produces the best sites in combination with a glutamate residue in position $i - 4$.³³⁰ As secondary structure predictions identify amphipathic α helices with good reliability, this approach is of interest for proteins of unknown 3D structure.

3.3.4.3. Hybrid Double-Arm Attachment. Nitsche et al. recently introduced a new tagging strategy, where a small metal binding tag is attached to two cysteine residues, but additional coordination of the lanthanoid ion by a negatively charged side chain of the protein is still required to immobilize the metal ion.³³¹ The concept was demonstrated with the Zika virus NS2B-NS3 protease mutated to contain cysteine residues in positions 101 and 131. The side-chains of the cysteine residues are in close proximity and avidly bind the pnictogens arsenic,

antimony, and bismuth in oxidation state III. The third coordination site of the pnictogen is available for binding a mercaptomethylaryl tag, such as **4MMDPA** to capture a Ln(III) ion, which was shown to generate sizable $\Delta\chi$ tensors in conjunction with additional coordination by a glutamate residue. The fundamental advantage of the concept is its selectivity for two cysteines in close proximity. As single thiol groups bind pnictogens with negligible affinity, it allowed tagging of the protein in the presence of the native, solvent exposed single cysteine residues.

3.3.5. Cosolute Paramagnetic Probes. Cosolute paramagnetic NMR probes do not chemically modify the targeted biological macromolecule. A structurally diverse class of paramagnetic molecules has been designed and investigated for their noncovalent interaction with the solvent exposed surface residues of oligopeptides and proteins. Cosolute probes can be as simple as salts of paramagnetic transition metals. The approach is based on the idea that large compounds and charged

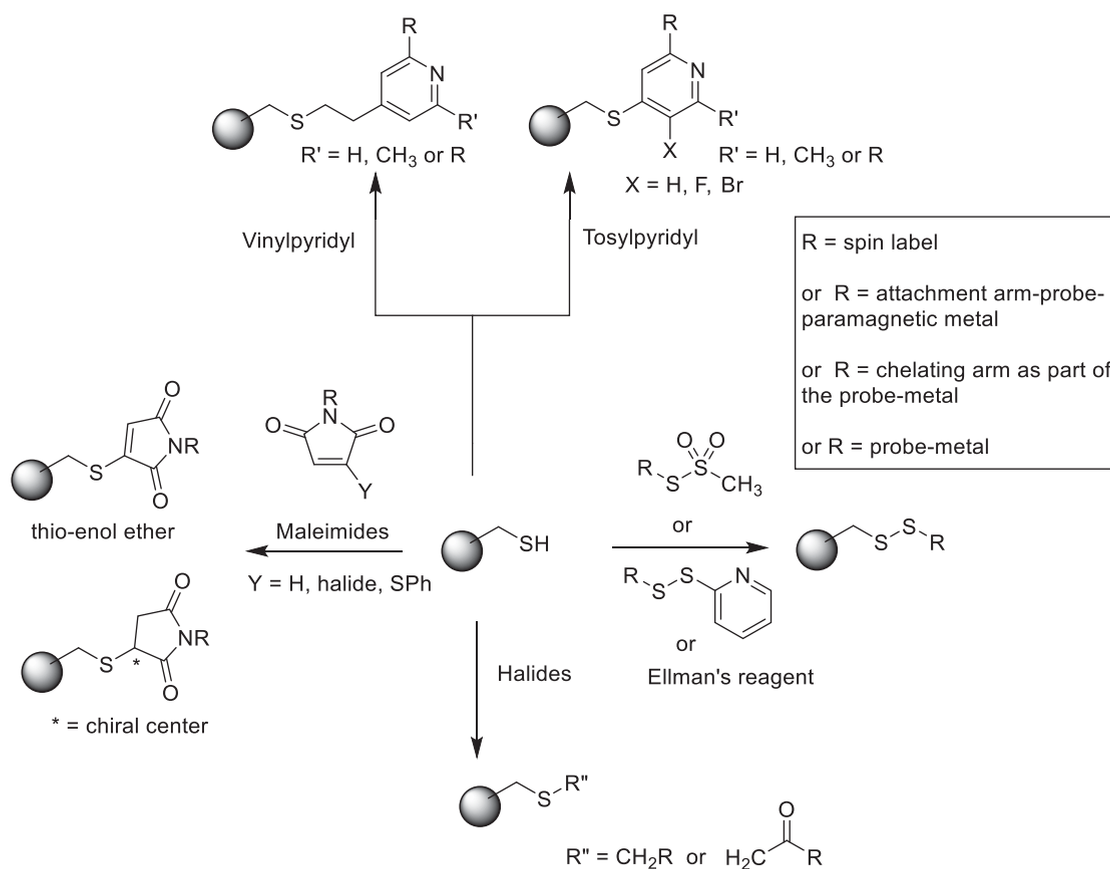


Figure 14. Most relevant chemical methods to connect NMR or EPR probes to one or more cysteine residues in proteins. The aim of the figure is to provide an overview of the currently used chemistry for the modification of cysteine residues, suitable for the attachment of a chemically sensitive paramagnetic center. The use of maleimide as reactive group results in the generation of a chiral center, and therefore, this type of linkage is not recommended in NMR studies.

ions do not readily access the interior of folded proteins, so that the NMR signals of surface residues experience stronger PREs than buried residues.³³² Mn(II) ions are well suited for this type of NMR experiments.^{333–335}

Owing to its diradical character, molecular oxygen is paramagnetic and one of the simplest molecular probes used in NMR. As oxygen dissolves more easily in the hydrophobic interior of membranes than in aqueous solution, this solubility gradient has been used to probe the structure and positioning of a membrane protein,³³⁶ and a number of related NMR experiments have been reported that exploit the unique properties of oxygen.^{337–341} In early work, Williams and co-workers obtained cosolute paramagnetic effects by directly adding paramagnetic metal ions, such as Gd(III), Eu(III), and Mn(II), to protein solutions.^{110,120,342} Using $[\text{Cr}(\text{CN})_6]^{3-}$, $[\text{Fe}(\text{CN})_6]^{3-}$, and $[\text{Co}(\text{CN})_6]^{3-}$ complexes in 1D ^1H NMR spectroscopy experiments, Williams and co-workers studied the competition with negatively charged substrate molecules for binding to phosphoglycerate kinase.¹¹¹

Organic nitroxide radicals present a more varied class of paramagnetic cosolute probes. Early results showed that di-*tert*-butylnitroxide generates contact shifts with several small organic molecules in carbon tetrachloride.³⁴³ On the basis of these results, Kopple and Schamper³⁴⁴ prepared 3-oxyl-2,2,4,4-tetramethylloxazolidine (Figure 12) that is soluble in chloroform and methanol, and these solvents and this probe were used to detect line broadening in the ^1H NMR spectrum of amides of the solvent exposed amino acid residues in the C_2 symmetric cyclic

decapeptide gramicidin S, while the amides of the amino acid residues involved in intramolecular H-bonding were not affected. The same research group reported that a water-soluble variant (HyTEMPO or TEMPOL, Figure 12) can serve as a tool to identify hydrophobic surface residues of proteins, using the attenuation of antiphase COSY cross-peaks as a sensitive method to detect small line-broadening effects caused by the presence of the paramagnetic agent.³⁴⁵ Building on these results, Fesik and co-workers showed that quantitative measurement of $R_1(^1\text{H})$ relaxation rate enhancements generated by TEMPOL allowed accurate identification of the solvent exposed regions of cyclosporin A bound to cyclophilin.³⁴⁶ TEMPOL also proved useful to probe the accessibility of amino acid residues in protein unfolding experiments³⁴⁷ and studies of conformational change.³⁴⁸ The attempt to probe electrostatic potentials using the charged derivatives 4-carboxy-TEMPO and 4-amino-TEMPO (Figure 12) showed the expected trends but also demonstrated limitations attributed to local variations in translational diffusion constants.^{347–349} Stronger solvent PREs can be generated by Gd(III) ions and the uncharged and highly water-soluble Gd(III)DTPA-BMA probe (Figure 13) can perform better than TEMPOL, because it is less hydrophobic, stable with regard to redox-active compounds, and the paramagnetic center less accessible.³⁵⁰ Solvent PREs generated by Gd(III)DTPA-BMA were shown to deliver powerful restraints for 3D structure determinations of proteins and peptides,^{35,351,352} which prompted the design of Gd(III)-TTHA-TMA^{353,354} as a more spherical Gd(III) probe without

any hydration water bound to the Gd(III) ion. The use of alternative uncharged Gd(III) probes such as $\text{Gd}_2(\text{L}_7)\text{-(H}_2\text{O)}_2$,³⁵⁵ $\text{Gd}(\text{DO3A})$, and $\text{Gd}(\text{HP-DO3A})$,³⁵⁶ as well as complexes with increasing charge, such as $[\text{Gd}(\text{EDTA})]^{1-}$,³⁵⁷ $[\text{Gd}(\text{DTPA})]^{2-}$,³⁵⁸ and $[\text{Gd}(\text{DOTP})]^{5-}$,³⁵⁹ has also been explored (Figure 13). The strong PRE effects generated by Gd(III) complexes have also proven useful to shorten the T_1 relaxation time of H_2O ³⁶⁰ and protein protons³⁶¹ without greatly increasing the line widths of protein NMR signals, enabling sensitivity enhancements by faster scan repetition rates. It is important to note, however, that some proteins may harbor specific binding sites for some of these complexes, which can be made apparent by PCSs generated by complexes with paramagnetic lanthanoid ions other than Gd(III). For example, the cosolute probe $[\text{Tb}(\text{DOTP})]^{5-}$ was found to generate specific PCSs in the DNA binding protein fd gene 5³⁵⁹ and $[\text{Ln}(\text{DPA})_3]^{3-}$ (with Ln = Tm, Yb, or Tb, Figure 13) was found to generate significant PCSs in ArgN and several other proteins.^{362,363} The $\Delta\chi$ tensor generated in this way can be tuned by using DPA derivatives carrying different substituents ($-\text{Br}$, $-\text{COOH}$, or $-\text{CH}_2\text{OH}$) on the aromatic ring, which form lanthanoid complexes like the parent $[\text{Ln}(\text{DPA})_3]^{3-}$ and bind to the same site on the protein.³⁶³ As rapid reorientation of the complexes averages PCSs to zero,³⁵⁰ the observation of PCSs can be taken as positive proof of specific binding. Natural binding sites for $[\text{Ln}(\text{DPA})_3]^{3-}$ are infrequent in proteins and it has been shown that binding sites specific for this complex can be engineered by positioning the side chains of two positively charged residues (lysine or arginine) in close proximity, which, at the same time, need to be sufficiently far from negatively charged carboxylates to prevent their engagement in salt bridges.³⁶⁴ In contrast to covalently bound paramagnetic probes, specific binding sites for complexes such as $[\text{Gd}(\text{DPA})_3]^{3-}$ offer the opportunity to measure PREs for nuclear spins very close to the binding site, as the PRE magnitude can be tuned by the amount of relaxation agent added.³⁶⁵ PREs generated by the cyclen complexes $\text{Gd}(\text{DO3A})$, $\text{Fe}(\text{DO3A})$, and $\text{Ni}(\text{DO2A})$ (Figure 13) have also been explored. The Gd(III) ion in $\text{Gd}(\text{DO3A})$ has two accessible coordination sites available and specifically targets carboxylates on protein surfaces.³⁶⁶ $\text{Fe}(\text{DO3A})$ has been used to speed up data acquisition by shortening the T_1 relaxation time of an intrinsically disordered protein.³⁶⁷ $\text{Ni}(\text{DO2A})$ has been reported to be particularly suitable for shortening the T_1 relaxation times of H_2O and protein signals without broadening them significantly.³⁶⁸ As the ligand fills all coordination sites of the Ni(II) ion, it is unlikely to bind to specific sites on proteins. At the same time, it is a water-soluble probe without carrying a net charge.

3.3.6. Development of Cysteine Based Probe Attachment Chemistry. Historically, the development of chemical approaches for the selective modification of proteins has its origin in studies of amino acid chemistry.³⁶⁹ Further important insights on site-specific chemical modification came from studies of post-translationally modified proteins.³⁷⁰ New chemistry for site-specific ligation reactions were opened by the development of methods to incorporate noncanonical amino acids either by genetic encoding or the use of chemically miscyclated tRNAs.^{371–373}

The thiol group of cysteine has unique nucleophilic characteristics, as compared to the side chains of any of the other canonical amino acid residues. Because of the higher acidity of RCH_2SH as compared to RCH_2OH , cysteine residues

in proteins can be deprotonated at slightly basic pH, which increases their reactivity^{374–383} toward soft electrophiles. With little chemical activation of the paramagnetic probes, the reaction conditions used for the attachment of the probes can be mild. Figure 14 provides an outline of the most relevant cysteine modifications with respect to the attachment of probes amenable for NMR and EPR.^{384–386}

A standard, selective and mild, method for attaching a probe to a cysteine residue is by disulfide formation. One of the most frequently used reactive functionalities to accomplish this is the MTS or S-mesyl group. This group is stable under acidic conditions, which is advantageous, if acid labile protective groups need to be removed at a late stage of chemical probe synthesis.²⁸⁰ At physiological pH the thiol of cysteine forms a disulfide linkage with the RS-moiety of the S-mesyl compound with the release of methane and SO_2 in a practically irreversible reaction. Several other reactive functionalities are known to generate S–S bonds,³⁸⁷ in particular the 2-pyridyldisulfanyl group (as in the probes 60,²⁴⁸ 82,²⁷⁸ 83,²⁷⁹ 86,²⁸² 94,¹²⁶ 96,²⁹⁰ 97,²⁹² 101,¹²⁹ 103,¹²⁹ 112,³⁸⁸ and 127³²⁹). Disulfide bonds are also made by activation of a cysteine residue of a protein using a mild oxidizing agent such as Ellman's reagent (DTNB) and addition of the probe as a thiol.^{322,323,327,328,389} Similarly, a chelating moiety was introduced by reaction of a cysteine residue with ArS-SAr ($\text{Ar} = \text{PyMTA}$ ²⁶⁶ or $\text{Ar} = \text{DPA}$ ³²⁴). Several procedures are available to produce a thioether, which is chemically more stable than a disulfide linkage. α -Iodo or α -bromo carbonyl reagents ($\text{XCH}_2\text{C}(=\text{O})\text{R}$) have traditionally been used to generate thioethers with cysteine residues, as the reaction is fast and selective at the physiological pH.^{369,390} This reactive functionality has successfully been installed in a number of probes, such as in probes 16,²¹³ 17,³⁹¹ 81,²⁷⁷ 87,²⁸⁵ and 104.²⁹⁶ Furnishing DOTA type probes with two α -bromo-carbonyl functionalities, however, has proven challenging because of the high reactivity of this functionality.²⁷⁷

Another standard reactive functionality is the maleimide group, which has a high reactivity and selectivity for cysteine residues around neutral pH.^{392,393} Maleimide probes react with cysteines via Michael addition, forming a new stereocenter^{211,212,281} and thus giving rise to different stereoisomers which complicate PCS assignments. The use of a maleimide functionality to attach a probe for NMR experiments is therefore not recommended. Monobromomaleimide probes^{223,394} do not have this problem because the Michael addition is followed by an elimination reaction resulting in a thio-enol ether linkage (Figure 14), and Michael reactions with vinylpyridine probes (probes 64,²⁵³ 65,²⁵³ 74,²⁶⁴ 75,²⁵ 121,³²⁶ and 123¹⁷⁶) do not generate new chiral centers either. Tosylpyridine probes (62,²⁵³ 63,²⁵³ 71,²⁶⁰ 72,²⁶⁰ 76,²⁶⁵ 77,²⁶⁶ 88,²⁸⁶ 89,²⁸⁷ 90,²⁸⁷ 91,²⁸⁷ 93,²⁸⁸ 105,²⁹⁷ 106,²⁹⁷ 107,²⁹⁸ 108,³⁰⁰ 109,³⁰¹ 110,³⁰¹ and 124²⁶⁵) form thioethers. The electrophilicity of these probes is enhanced by the presence of a halide substituent on the pyridyl ring. Similar to the tosyl group, a nitro group on the pyridine ring can act as a leaving group.²⁹⁴

3.4. Paramagnetic Probes for DNA and RNA

Studying oligonucleotides with paramagnetic probes can provide valuable insights into the structure and dynamics of DNA and RNA as well as their interactions with proteins.^{40–47} Labeling methods of oligonucleotides can be classified into four main categories. First, noncovalent spin labels can be selectively introduced at an abasic site without covalent attachment of the probe to the nucleotide. Second, spin labels can be selectively

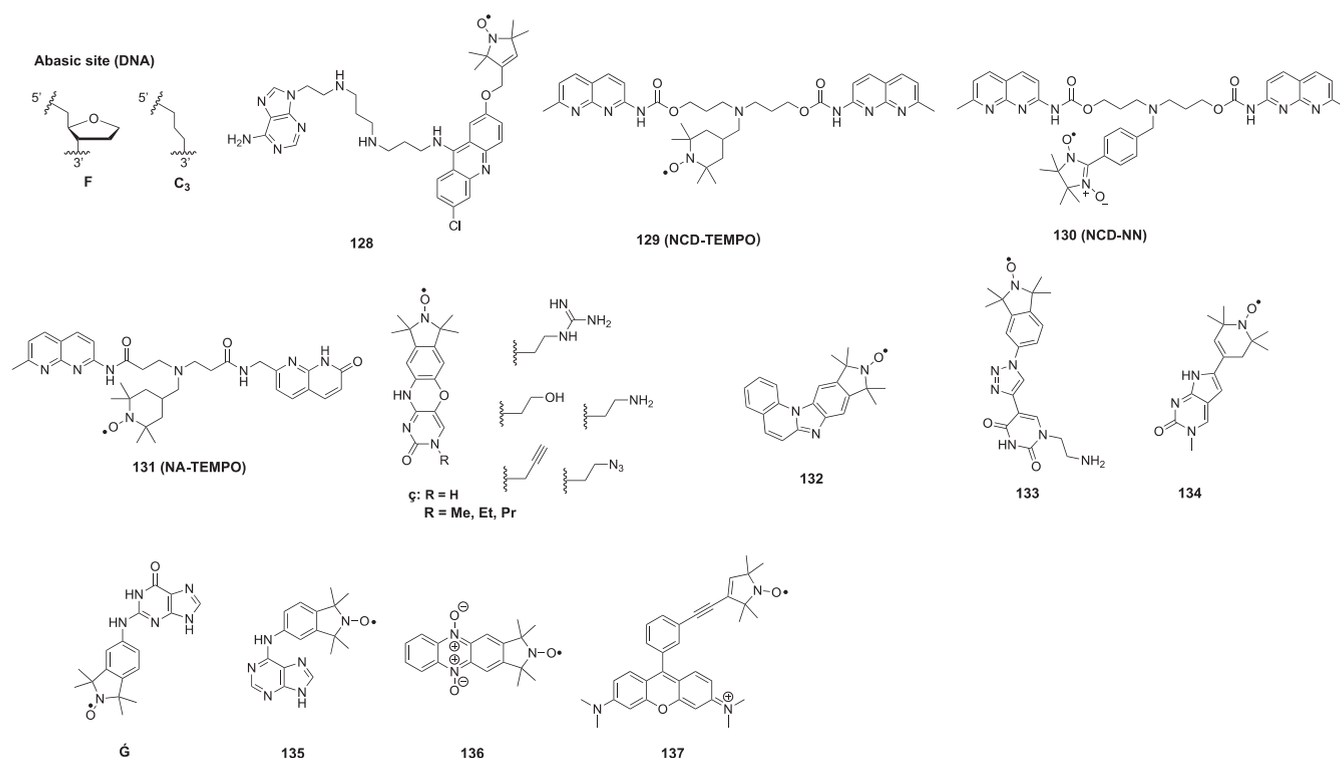
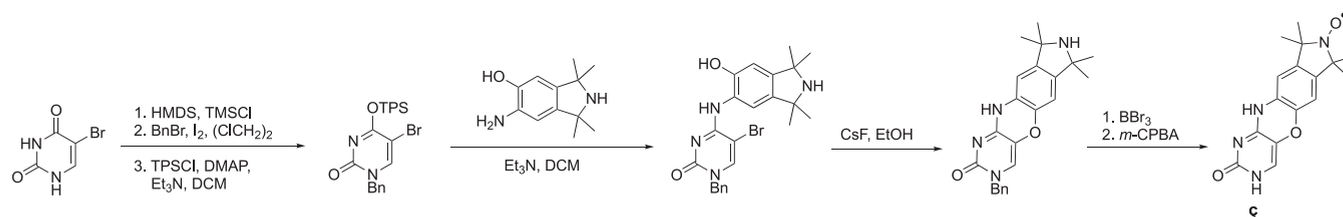


Figure 15. Site-directed noncovalent spin labels for DNA and RNA.

Scheme 9. Synthesis of the Noncovalent DNA Spin Label ϕ from 5-Bromouracil^a



^a5-Bromouracil is regioselective benzylated, subsequently converted into its O⁴-sulfonylated derivative (TPS = 2,4,6-triisopropylbenzenesulfonyl) and coupled to an isoindol aminophenol derivative, followed by ring-closure. The benzyl group is removed and N-oxidation gives the nitroxide.

attached to the nucleotide backbone via reaction with modified ribose or phosphate groups in the oligonucleotide. Third, spin labels can be selectively attached to a single base via reaction with a modified purine or pyrimidine base. Fourth, a fully synthetic and already labeled nucleotide can be introduced into oligonucleotides by standard solid-phase oligonucleotide synthesis. Occasionally, a synthetic method may allow the attachment of probes to the nucleotide prior to, during, or post oligonucleotide synthesis. All four strategies are reviewed in the following sections with a particular emphasis on the attachment chemistries.³⁹⁵

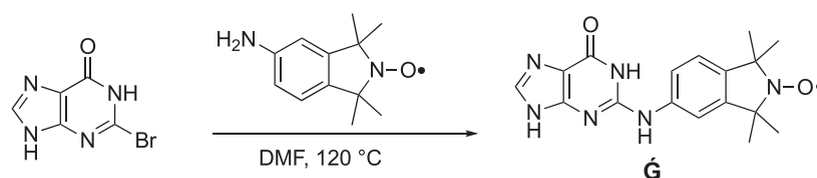
3.4.1. Noncovalent Spin Labels. Paramagnetic cosolutes, which have been used intensively for the study of proteins (section 3.3.5), have also been used to study oligonucleotides by paramagnetic NMR spectroscopy. For example, the nitroxide radical TEMPOL (Figure 12) and the Gd(III) complex of TTHA-TMA (Figure 13) have been used to study the surface accessibility of RNA and refine its 3D structure.^{396,397}

To probe biomolecules more precisely, it is necessary to introduce the spin label at a defined position as rigidly as possible. In the case of proteins, this is usually achieved by site-selective mutation and subsequent covalent bond formation

between probe and amino acid side chains (section 3.3). Selective and rigid noncovalent labeling of proteins is not straightforward and achievable only in special cases, requiring the availability of a suitably labeled protein ligand.³⁹⁸ In the case of oligonucleotides, noncovalent spin labeling is a much more generally applicable technique that appeals by its simplicity. Synthetic spin labels can mimic a purine or pyrimidine base and specifically bind to an abasic site in double-stranded DNA or RNA oligomers, thus circumventing the need to synthesize the entire spin-labeled nucleotides. Currently available noncovalent spin labels are all nitroxide radicals, which allow applications in NMR (PRE) and EPR. Noncovalent oligonucleotide tags containing a paramagnetic metal ion may be suitable for PCS measurements but are yet to be explored.

The first site-directed noncovalent spin label, which binds to an abasic site in DNA, was reported by Lhomme and co-workers (Figure 15, 128).^{399,400} Compound 128 combines the base adenine with the DNA intercalator acridine, which are connected via a flexible linker. Nakatani and co-workers developed spin label NCD-TEMPO (Figure 15, 129) and a nitronyl nitroxide analogue NCD-NN (Figure 15, 130), which can specifically bind to G–G mismatches in DNA duplexes,

Scheme 10. Synthesis of the Noncovalent RNA Spin label \dot{G} from 2-Bromohypoxanthine via a Nucleophilic Aromatic Substitution



allowing for the construction of tandem arrays and three-dimensional assemblies of electron spins.^{401–404} Spin label NA-TEMPO (Figure 15, 131) was designed in this work to clarify the effect of orthogonal sequence selectivity on the electron-spin arrangement.⁴⁰²

Sigurdsson and co-workers introduced the noncovalent DNA spin label ζ , which allows labeling without intercalation or specific sequence requirements.⁴⁰⁵ Its synthesis from 5-bromouracil is shown in Scheme 9. Mimicking a cytosine, ζ binds best to a DNA duplex if paired with a guanosine nucleotide ($K_d \sim 5 \mu\text{M}$ at -10°C).⁴⁰⁵ Binding of ζ (Figure 15) does not only depend on the pairing base, but also on the flanking sequence, with a 5'-d(D ζ T) sequence affording the highest affinity.⁴⁰⁶ The abasic site should also be located at least four base pairs from the end of the DNA duplex.⁴⁰⁶ A DNA duplex labeled with two ζ labels enabled DEER distance measurements up to 70 Å. Binding of a double-stranded *lac* operator DNA sequence to the *lac* repressor protein has also been probed by DEER. Bending of the double helix was manifested in a distance decrease between both ζ spin labels by about 5 Å, which, however, is close to the resolution limit of this method.⁴⁰⁷ The introduction of substituents at the N3 nitrogen of ζ affected the affinity, with large substituents compromising the affinity most.^{406,408} Derivatives containing an amino or guanidino group displayed slightly increased affinity and improved solubility.⁴⁰⁸ The nucleoside C₃ (Figure 15) was explored as an alternative abasic site over the tetrahydrofuran analogue F (Figure 15) with limited impact on the affinity of ζ to the DNA.⁴⁰⁶

The nitroxide radical 132 (Figure 15) was developed as an alternative noncovalent spin label that binds to abasic sites in RNA and DNA duplexes.⁴⁰⁹ Best pairing was observed with cytosine which does not interact well with ζ . Although spin labeling of RNA and DNA was confirmed by EPR spectroscopy, 132 has not been used in any further applications. Another campaign screened ten pyrimidine based spin labels for their suitability to occupy abasic sites in DNA and RNA duplexes.⁴¹⁰ Triazole linked (e.g., 133 (Figure 15)) and pyrrolocytosine-based (Figure 15, 134) labels showed very promising results. Spin label 133 displayed increased solubility and affinity (DNA) compared to its N1-methylated analogue (not shown). It was speculated that the increased affinity may result from a salt bridge formed between the ethylamino group and the phosphate backbone. The DNA label ζ has insufficient affinity to RNA ($\sim 30\%$).⁴¹⁰ In contrast, 133 shows much higher affinity (100% under identical conditions) to an abasic RNA site; it also binds to an unmodified RNA duplex, which, however, compromises specificity.⁴¹⁰

The isoindoline-nitroxide derivative \dot{G} overcomes previous limitations and binds with high affinity and specificity to abasic sites in RNA duplexes. \dot{G} pairs with a cytosine and is suitable for DEER distance measurements when combined with an RNA duplex containing two abasic sites.⁴¹¹ \dot{G} can be prepared from

commercially available materials in a single straightforward synthetic step (Scheme 10), facilitating its broad use.⁴¹¹ \dot{G} can also bind to abasic sites in DNA-RNA hybrids, either in the RNA or DNA strand, and, similar to ζ , must be positioned at least four residues from the end of the duplex to achieve optimal binding.⁴¹² The adenine derived spin label 135 complements the guanidine derived \dot{G} (Figure 15); 135 was identified from a series of purine derived nitroxides and can be introduced at abasic sites in DNA and RNA duplexes where it pairs with thymine or uracil, respectively.⁴¹² \dot{G} has been successfully applied to measure distances in both DNA and RNA duplexes by DEER.⁴¹³ Combined with molecular dynamics simulations, these experiments indicate strong Watson–Crick base pairing, no disturbance of the overall helical structure (minor local disturbances occur) and rigid binding of \dot{G} . Recently, \dot{G} was also used to label DNA and RNA triplexes containing a single abasic site.⁴¹⁴ The *N*-oxide phenazine 136 is the latest noncovalent spin label that can bind to an abasic site in DNA but is unsuitable for RNA.⁴¹⁵

If the spin label can directly bind to RNA or DNA, no chemical modifications of oligonucleotides (not even the introduction of an abasic site) are required. Despite the principal appeal of this strategy, however, achieving the necessary specificity of the interactions between label and oligonucleotide can be challenging, as observed for intercalating spin labels based on chlorpromazine,⁴¹⁶ ethidium bromide, or other polyaromatic carcinogens,⁴¹⁷ Ru(II)-phenanthroline complexes,⁴¹⁸ or acridine.⁴¹⁹ Bifunctional cross-linking agents are also of limited specificity.^{420,421} One solution is presented by the use of an aptamer. For example, the malachite green RNA aptamer (a 38-nucleotide sequence) is known to bind malachite green or closely related dyes such as tetramethylrosamine (TMR). If TMR is synthetically combined with a spin label as in 137 (Figure 15), it can be used to selectively label the RNA aptamer.⁴²² Combination with a spin label that is covalently attached to the RNA backbone (145, see section below) allowed DEER measurements. Similarly, a ternary complex consisting of a DNA oligomer, the molecule chromomycin-A₃ and paramagnetic Co(II), where Co(II) was bound to two oxygens of the chromophore, has been used to measure PCSs and improve the structure of the complex.^{423,424}

3.4.2. Probe Attachment to the Sugar–Phosphate Backbone. Some paramagnetic spin labels can be attached to the phosphate backbone of DNA and RNA. It has been noted that spin labels attached to phosphodiester interfere less with duplex formation than other labels, as they are located at the edges of the polynucleotide helices.⁴³ The greatest disadvantage of this approach is that labeling of phosphodiester inevitably results in the formation of diastereomers, which need to be separated by chromatography and identified, if unambiguous structural information is to be obtained from paramagnetic NMR experiments or EPR studies.^{296,425,426} Recently, a synthetic strategy for the chiral synthesis of phosphorothioates

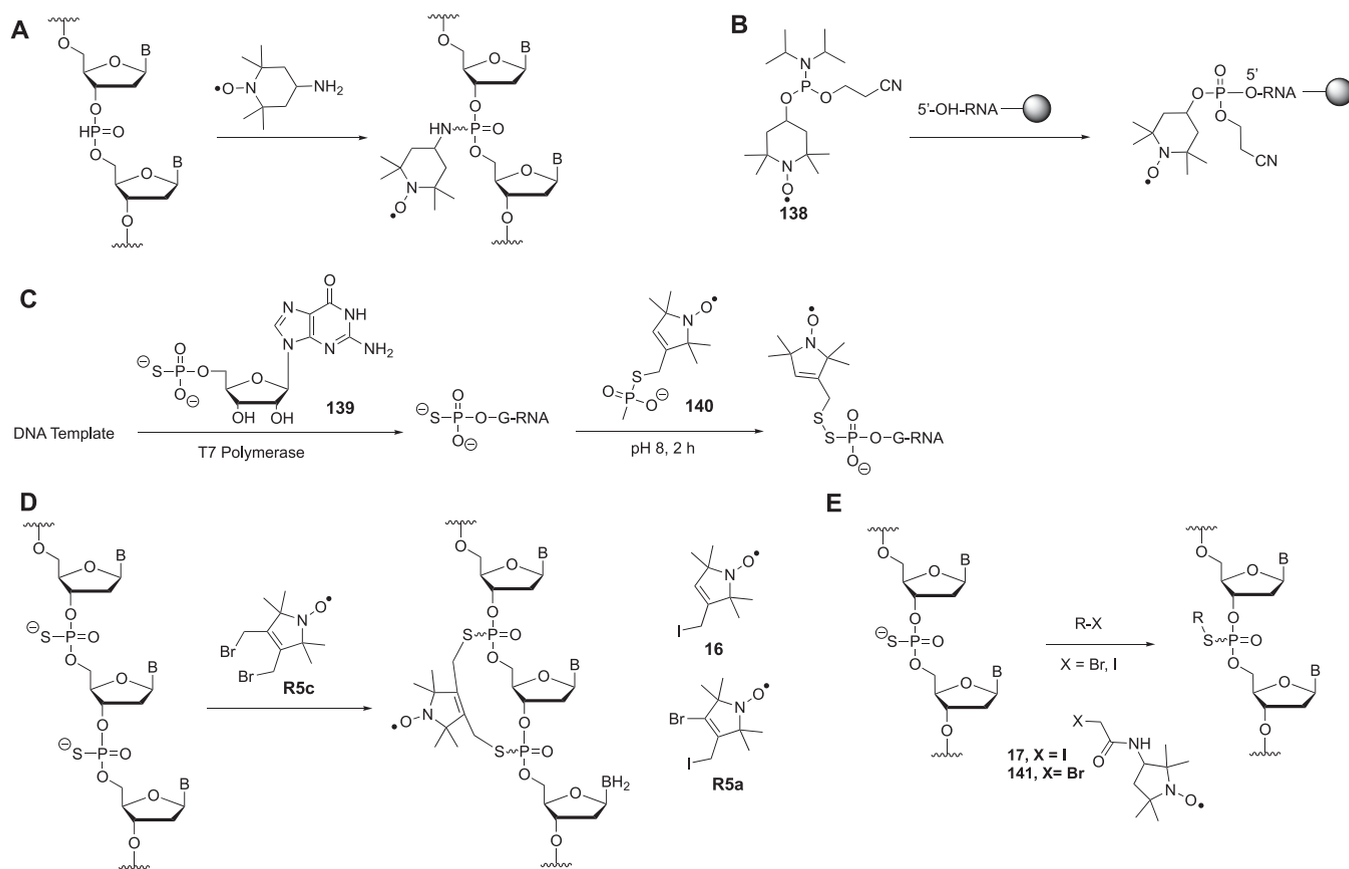


Figure 16. Strategies for the attachment of probes to the phosphodiester backbone or terminal phosphate groups of oligonucleotides.

has been described, which may help to overcome these challenges.⁴²⁷ In addition, it is important to note that, when labeling RNA, the 2' hydroxyl group adjacent to the labeled phosphodiester needs to be protected as a 2'-OMe group or replaced by a hydrogen atom to prevent strand cleavage through 2',3'-transesterification.⁴²⁸

The introduction of a hydrogen phosphonate group during oligonucleotide synthesis enables the selective attachment of **4-amino-TEMPO** via H-phosphonate chemistry yielding a phosphoramidate (Figure 16A).^{429,430} Using protected oligonucleotides, 4-hydroxy-TEMPO can be attached to the 3' or 5' end of DNA via a phosphoester bond (not shown),⁴³¹ which has been used to study protein DNA interactions by NMR.⁴³² Similarly, the increased reactivity of terminal phosphate groups can be exploited by selective activation of the 3' and 5' ends of unprotected DNA and subsequent reaction with **4-amino-TEMPO** to form phosphoramidates.⁴³³ A fully protected TEMPO-phosphoramidite building block (Figure 16B, **138**) compatible with oligomer synthesis has been used to form an ester between **4-hydroxy-TEMPO** and the phosphate at the 5' end of RNA.⁴³⁴

A frequently used functional group for site-directed labeling of the phosphate backbone is presented by phosphorothioate. Selective incorporation of spin labels at the 5' end of RNA is possible directly by transcription from DNA in a two-step process where, first, a DNA template is transcribed in the presence of NTPs (nucleotidetriphosphates) and a phosphorothioate nucleotide and, second, a spin label is attached to the 5' phosphorothioate group. The approach has been demonstrated with guanosine monophosphorothioate (**139**), T7 RNA

polymerase, and spin label **140**, which was attached via disulfide formation (Figure 16C).⁴³⁵

Alternatively, a phosphorothioate moiety can be enzymatically installed in the 5' position of RNA or DNA using T4 polynucleotide kinase, prior to alkylation with the spin label **16** (Figure 16).⁴³⁶ The iodoacetamide based spin label **17** (Figure 16) can be ligated to internal phosphorothioate nucleotides in DNA (Figure 16E).⁴³⁷ The rigid spin label **16** has been used more frequently.⁴³⁸ It has been attached to RNA and DNA to study, for example, the GAAA tetraloop/receptor interaction,⁴³⁹ the human telomeric G-quadruplex,⁴⁴⁰ riboswitch dynamics,⁴⁴¹ CRISPR-Cas9,⁴⁴² DNA structure and dynamics,⁴⁴³ distances by DEER,⁴⁴⁴ and protein DNA interactions.⁴⁴⁵ The 4-bromo-substituted analogue **R5a** has been used to study ribozymes, CRISPR-Cas9, and the structure and dynamics of DNA.^{426,446–449} The double-alkylating analogue **R5c** links two adjacent phosphorothioates to give a conformationally highly constrained cyclic spin label for RNA and DNA (Figure 16D).²²⁰ Graham and co-workers developed the first lanthanide tag (Figure 9, **C10**) that can be attached to phosphorothioate, enabling the observation of PCSs in NMR spectra of DNA.²⁹⁶

Alternatively, paramagnetic spin labels can be attached to the sugar (ribose) part of the backbone. The 2' position in oligonucleotides is the only position readily available for modification. Nucleotides bearing a 2' amino group can be site-specifically installed in oligonucleotides using standard oligonucleotide synthesis (2' amino-modified phosphoramidites of pyrimidine-based nucleotides are commercially available).^{450,451} The most commonly used spin label is **4-isocyanato-TEMPO** (Figure 17, **142**), which is readily accessible from **4-amino-TEMPO** and diphosgene.^{450,452} The

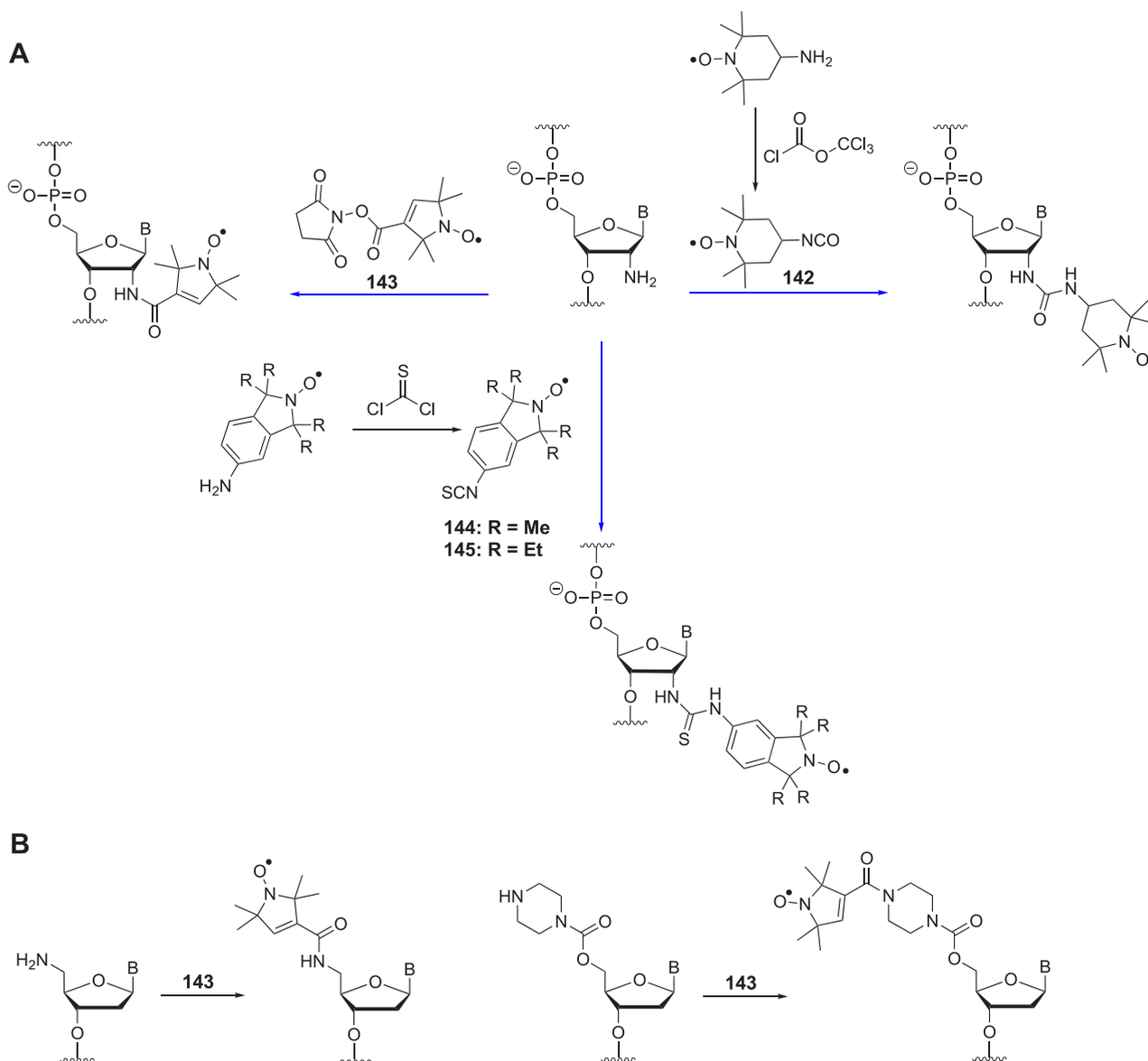


Figure 17. Probe attachment to (A) a 2' amino group of internal nucleotides (blue arrows indicate the labeling reactions) and (B) alternative 5' modifications.

isocyanate group forms a stable and rigid urea moiety between the 2' position of the nucleotide and the TEMPO spin label, which has been used in numerous applications involving EPR and NMR to study RNA and its interactions.^{453–461} An alternative succinimide-based spin label (Figure 17, 143) was found to affect the thermostability of RNA helices.⁴⁶² It can also be introduced at a 5' amine or piperazine group (Figure 17).⁴⁶³ The isoindoline-based spin labels 144 (Figure 17) and 145 (Figure 17) bear an isothiocyanate group to form a thiourea linkage with the 2' amino group.²³⁰ The tetraethyl derivative 145 (Figure 17) displayed significant stability toward reducing conditions.

Attaching a probe to the 2' position via “click” chemistry has also been studied extensively (Figure 18). Attachment of probe 24 (Figure 18) to a 2' propargyl ether was the first example described; however, the construct produced broad distance distributions in DEER experiments due to a rather long and flexible linker.^{43,464} A much shorter linker can be established, if the alkynyl group is directly connected to the 2' carbon as in 2'-

alkynyl nucleotides (Figure 18).^{225,465} Azide based spin labels 23–25 (Figure 18) and 146–148 (Figure 18) can be attached selectively via “click” chemistry and have been used to measure distances in DNA by DEER.²²⁵ A crystal structure of double-stranded DNA with label 24 has been solved (Figure 19).⁴⁶⁵ Combined with EPR experiments and MD simulations, the results indicate only minimal perturbation of the DNA structure because of the presence of the spin label. Royzen and co-workers developed alkyne substituted Cu(II) (Figure 18, 149) and Co(II) (Figure 18, 150) binding ligands that can be attached to a linker-azide functional group in the 2' position (Figure 18).^{466,467} Both tags were successfully used to study binding of the HIV-1 nucleocapsid protein 7 to an RNA pentanucleotide by PRE measurements using NMR spectroscopy.^{466,467} In the case of the cobalt tag 150, observation of PCSs was not reported, which potentially relates to the flexibility of the linker between probe and RNA.⁴⁶⁷ The “click” reaction between a 2' azido GTP and an alkyne-TEMPO derivative produces a spin labeled nucleotide that can be used for site-specific post-transcriptional

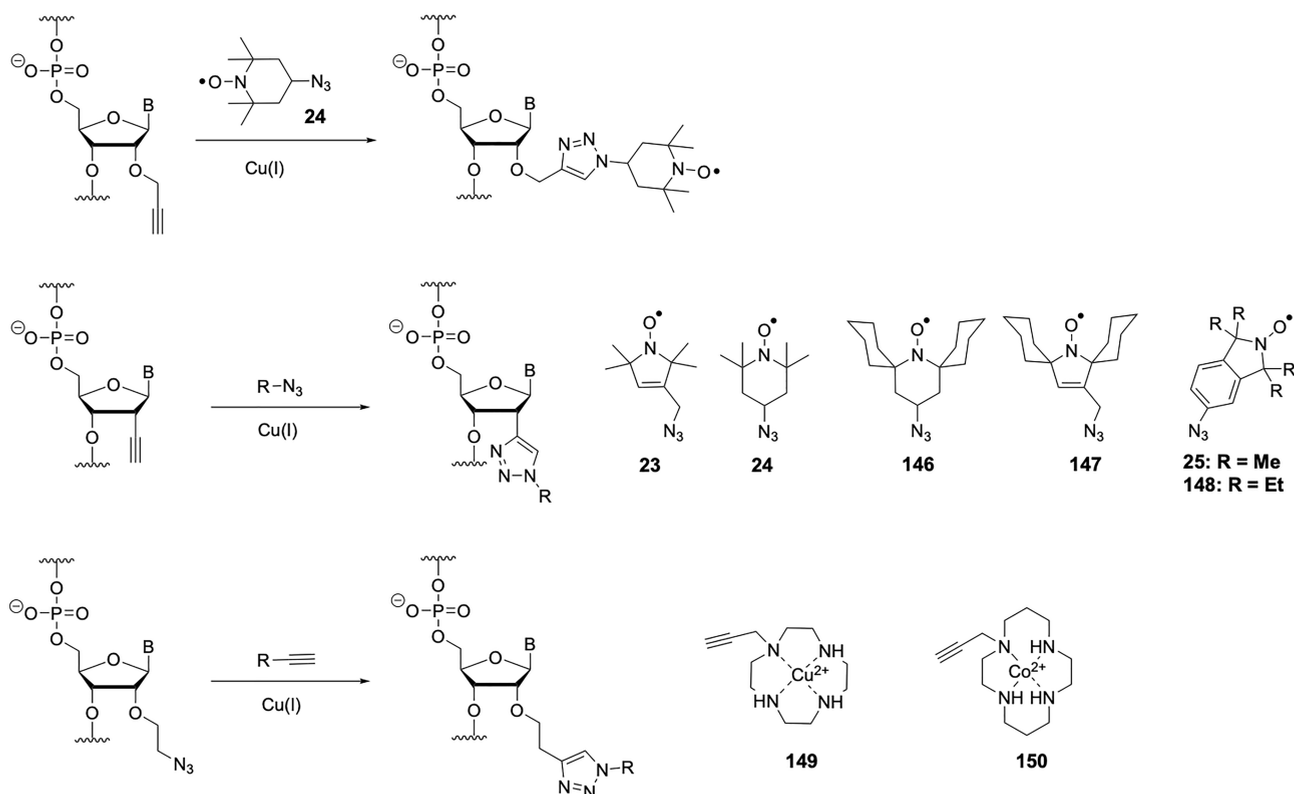


Figure 18. Probe attachment via “click” chemistry to the 2' position.

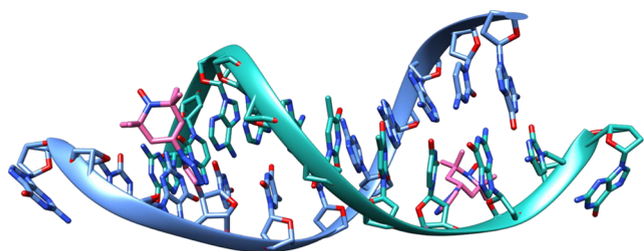


Figure 19. X-ray crystal structure (PDB code 6QJS) of a 12 base pair DNA duplex containing the spin label **24** (highlighted in pink) attached in the 2' position via “click” chemistry. The phosphate backbone is shown as a ribbon. The distance between both nitroxide radicals determined by DEER is 30 Å.

labeling of RNA (not shown).⁴⁶⁸ The GTP analogue is a substrate for a Tb^{3+} -deoxyribozyme that installs it specifically on an internal adenine nucleotide via a 2',5'-phosphodiester linkage.⁴⁶⁸

Finally, spin labels can be selectively attached to the 5' or 3' hydroxy ends of nucleosides. Selective 5' labeling can be achieved via carbamate or amide formation with TEMPO,⁴³⁰ **143** (Figure 17B),⁴⁶³ or the large trityl radical **151** (TAM) (Figure 20).⁴⁶⁹ Attachment to both 5' nucleotides of a DNA duplex allows distance measurements by DEER. Although trityl radicals offer advantages over nitroxide radicals, such as narrow spectral width and stability under reducing conditions,⁴³ their size and hydrophobicity limit applications with biomolecules. Recently, alternative linkers were developed that allow the attachment of TAM anywhere in the DNA sequence using an achiral non-nucleoside phosphoramidite (Figure 20).⁴⁷⁰ Selective modification of the 3' end of RNAs with TEMPO has also been demonstrated.^{471–473}

3.4.3. Probe Attachment to Modified Bases. The majority of paramagnetic probes used to study RNA and DNA are attached to purine or pyrimidine bases. Labels can be fully introduced by synthetic chemistry to furnish labeled nucleotides that can subsequently be used directly in solid-phase oligonucleotide synthesis (covered in the next section). Alternatively, labels can be introduced postsynthetically (by chemical synthesis or transcription) by attachment to a modified base either prior or post removal of protection groups (in the case of chemical synthesis). The postsynthetic approach can help minimize possible decomposition of the nitroxide radical during synthesis, but it may result in incomplete labeling or side reactions with other functional groups.⁴³

A functional group frequently employed for postsynthetic modification is presented by sulfur substituents in purine and pyrimidine bases (Figure 21). 4-Thiouracil has been used most frequently. 4-Thiouridine can be found in natural tRNAs of *E. coli* and has been selectively labeled with spin label **17/141** (Figure 16) without the need for any synthetic modifications.⁴⁷⁴

Subsequently, 4-thiouridine was installed in synthetic RNA and labeled with **17/141** to study a double-stranded RNA-binding domain in the *Staufen* protein by NMR spectroscopy.⁴⁷⁵ A similar protocol⁴⁷⁶ was also used to determine the solution structure of large RNAs and protein RNA complexes,⁴⁷⁷ for example, to study *E. coli* protein sequestration by the noncoding RNA RsmZ by NMR and EPR.⁴⁷⁸ A very similar system using deoxy-4-thiouridine and **17/141** in synthetic DNAs (installed at a 5' overhang) allowed the study of DNA and the modulator recognition factor 2 (Mrf2) by NMR measurements of PREs.⁴⁷⁹ As an alternative to the formation of thioethers, thiouridine can form disulfide bonds with spin labels **152–154** (Figure 21), which has been used to study RNA structures.⁴⁸⁰ 2-Thiocytidine has been incorporated into tRNA using tRNA nucleotidyl

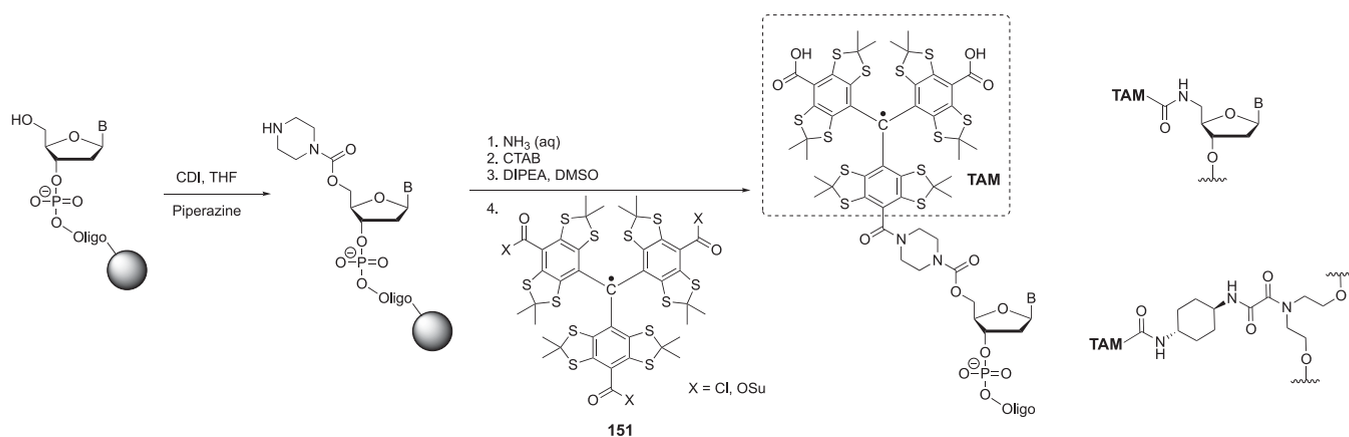


Figure 20. Attachment of the triarylmethyl (TAM)-based spin label **151** to the 5' end of DNA. The immobilized oligonucleotide is treated with CDI and piperazine and cleaved from the support using standard conditions. The resulting oligonucleotide is treated with cetyltrimethylammonium bromide (CTAB), taken up in DMSO and treated with **151** as the chloride in the presence of base. The two remaining acid chloride functions were hydrolyzed. Two alternative TAM attachments strategies are shown.

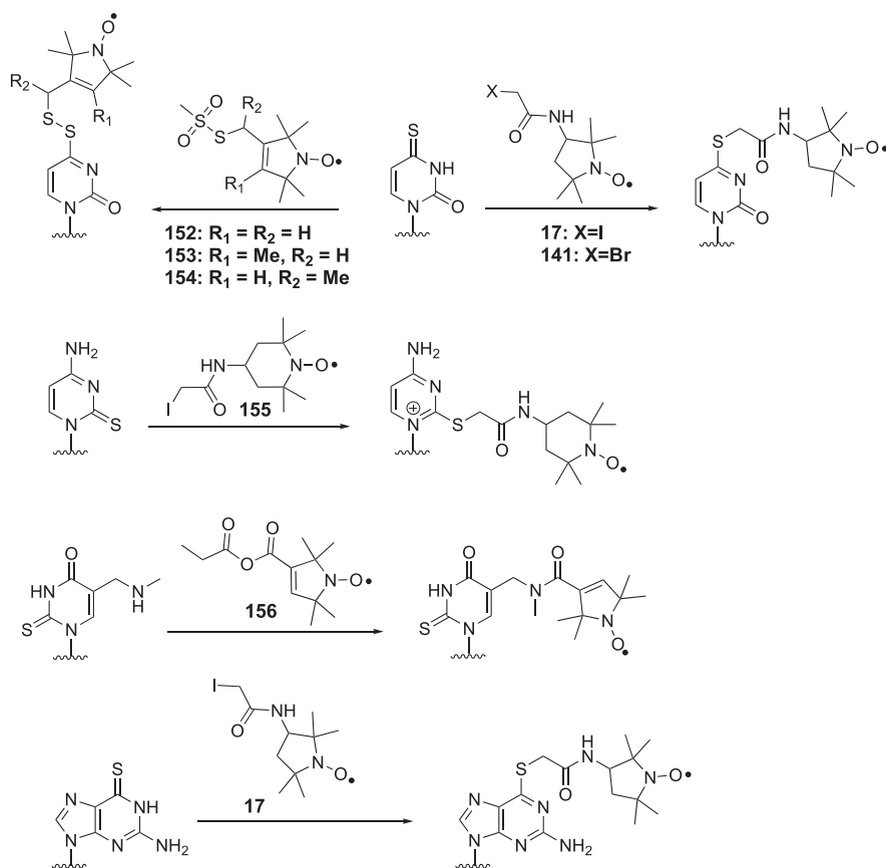


Figure 21. Postsynthetic probe attachment to thiopyrimidines and thiopurines.

transferase and reacted with spin label **155**.⁴⁸¹ The rare nucleotide 2-thio-5-(*N*-methylaminomethyl)-uridine from the anticodon region of *E. coli* tRNA^{Glu} was acylated with spin label **156**.^{482,483} The thiopurine nucleotide 6-thioguanosine can be enzymatically incorporated into RNA via a DNA-splint mediated ligation strategy and subsequently reacted with spin label **17** to study long RNAs by NMR spectroscopy (Figure 21).^{484,485}

Convertible nucleotides present yet another strategy to introduce a functional group during oligonucleotide synthesis that can selectively react with spin labels postsynthetically. This

strategy uses aromatic ethers or fluorine as leaving groups and amino groups in spin labels, such as 4-amino-TEMPO, as nucleophiles (Figure 22).^{486–489} Both pyrimidine and purine analogues have been described, and the approach has been employed in several studies involving RNA and DNA.^{490–495}

Alkynyl bases, such as the commonly used 5-ethynyl-2'-deoxyuridine (EdU), have been developed to allow for the selective reaction with azido based spin labels (e.g., **24**, **25**, **148**) via “click” chemistry during postsynthetic modification of DNA and RNA (Figure 23A).^{496–503} The tags have been demonstrated to be particularly useful for distance measurements by

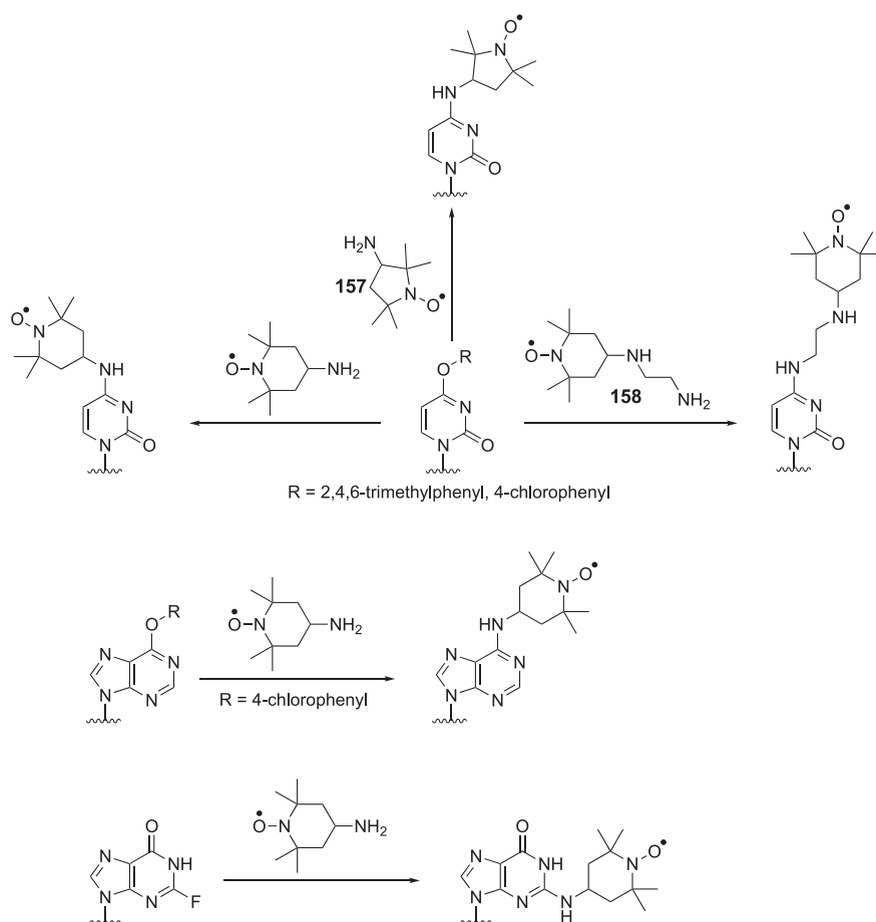


Figure 22. Convertible nucleotides for postsynthetic transformation into spin labels. The 2-fluoro-hypoxanthine derivative has also been used as O-NPE protected version.

DEER.^{496,497,499,500,502,503} The strategy also enables the labeling of very long RNAs (>400 nucleotides) and distance measurements within them.⁵⁰⁰ NHS esters **143** and **159** (Figure 23B) have been used to acetylate nucleotides that contain aliphatic amines (Figure 23B).^{504–507} Selective labeling of guanine involves a multistep process including the use of a complementary DNA reagent. Despite the length of the resulting linkers, the strategy successfully enabled DEER distance measurements in RNA duplexes. An analogue of cisplatin (**160**, Figure 23C) has also been used to deliver spin labels to DNA by ligand exchange with guanine (Figure 23C),⁵⁰⁸ enabling paramagnetic NMR measurements (PRE) that confirmed bending of the DNA duplex induced by the presence of the platinum complex.⁵⁰⁹ Using an on-column strategy, spin label **27** (Figure 23D) was coupled to iodo-nucleotides in RNA by the Sonogashira reaction.⁵¹⁰ The approach for 2-iodo-adenine is shown in Figure 23. Nucleotide modification via Diels–Alder reactions was also successfully investigated but not applied for EPR or NMR spectroscopy of DNA or RNA.⁵¹¹

The selective posttranscriptional labeling of RNA can be achieved by using an expanded genetic alphabet based on the unnatural base pair dTPT3-dNaM.⁵¹² Subsequently, the cyclopropene residue in the unnatural RNA nucleotide can be selectively modified with tetrazine reagents via a [4 + 2] cycloaddition with nitrogen release. If the tetrazine is linked to a spin label, as in **161** (Figure 23E), this approach can be used to introduce a paramagnetic tag into RNA after transcription from DNA.⁵¹³ Despite the very long linker, a distance in a self-

complementary RNA duplex could be determined by DEER. Alternatively, the dTPT3-dNAM system can be used to directly incorporate the spin-labeled nucleotide TPT3^{NO} into RNA by *in vitro* transcription, resulting in a much shorter linker and without the need for ligation chemistry (Figure 23E).⁵¹⁴

Different from proteins, lanthanoid tags for oligonucleotides are scarce. One reason is the catalytic effect of lanthanoid ions on hydrolytic cleavage of the phosphodiester bond if the metal ion is not fully protected by the complexation agent. Consequently, cyclen based tags, such as **C10**, are the preferred option.²⁹⁶ Recently, two TAHA based lanthanide tags were reported, which allowed the measurements of PCSs and RDCs in NMR spectra of double-stranded DNA.⁵¹⁵ Both tags **68** (Figure 7) and **162** (Figure 23F) can be attached via disulfide bond formation to a thiophenyl substituent in a selectively incorporated nucleotide derived from EdU (Figure 23F).

3.4.5. Fully Synthetic Paramagnetic Nucleotide Probes. Fully synthetic nucleotides are usually synthesized as their phosphoramidite analogues for subsequent use in standard solid-phase oligonucleotide synthesis. A summary of paramagnetic derivatives is given in Figure 24. TEMPO-labeled derivatives of 2'-deoxycytidine (Figure 24, **163**), 5-methyl-2'-deoxycytidine (Figure 24, **164**), 2'-deoxyadenosine (Figure 24, **165**), and 2-amino-2'-deoxyadenosine (Figure 24, **166**) were used in automated oligonucleotide synthesis, generating EPR-active DNAs.^{516–518} Because of the covalent link of the TEMPO group with the base, its mobility serves as a sensitive probe of the microenvironment, which was employed to detect

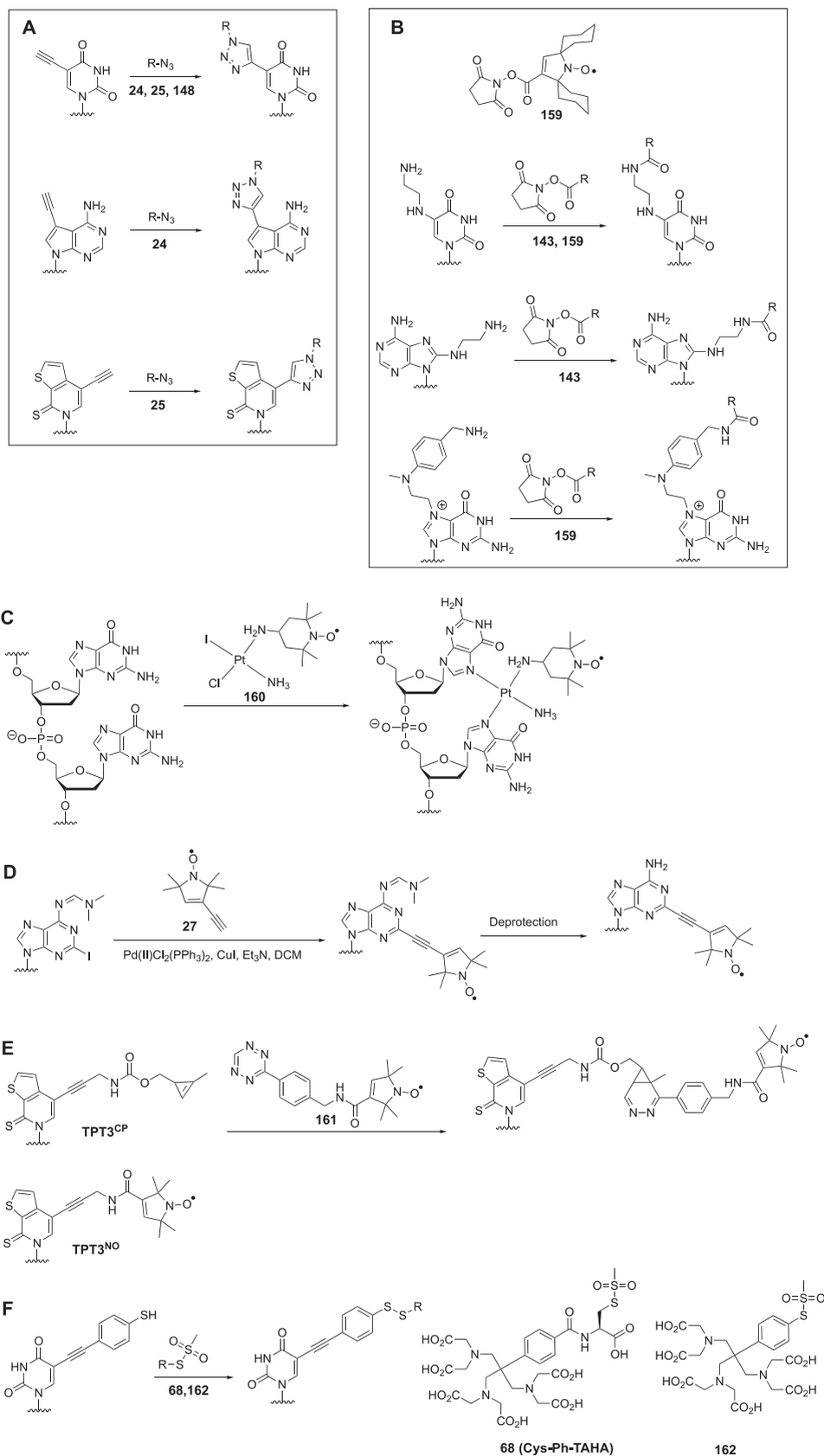


Figure 23. Various strategies for postsynthetic probe attachment to nucleotide bases.

single-base mismatches in duplex DNA by EPR using nucleotide 163.⁵¹⁹ TEMPO has also been attached via a semiflexible urea linker in analogues 167 (Figure 24) and 168 (Figure 24).⁵²⁰

The C5 position of pyrimidines is an alternative site for spin labeling that has been studied and applied intensively (Figure 24). Using phosphotriester chemistry, TEMPO nucleotides

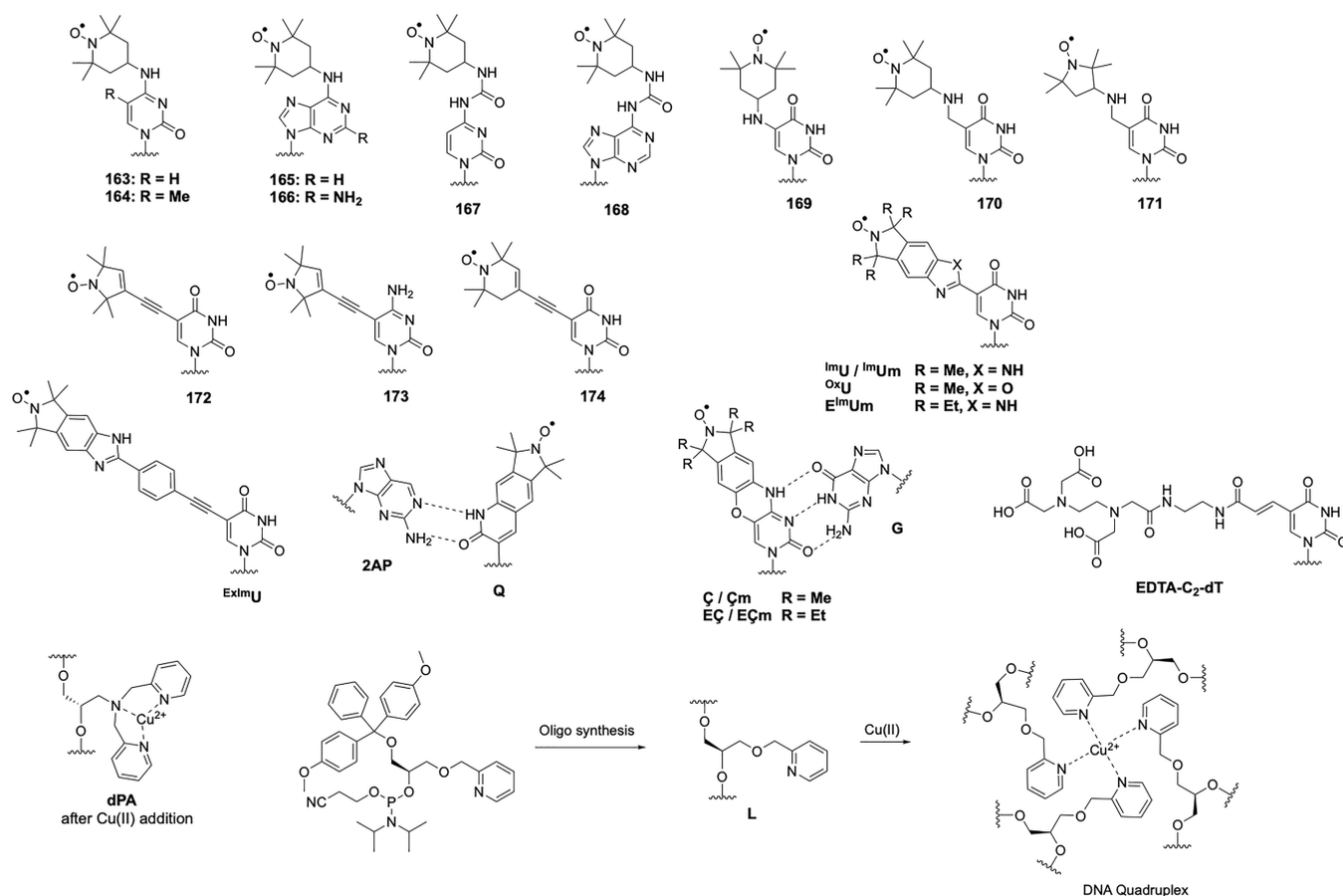


Figure 24. Paramagnetic nucleotides for incorporation during oligonucleotide synthesis.

(Figure 24, 169 and 170) and probe 171 (Figure 24) were incorporated into short DNA oligomers.^{521,522} Longer linkers between pyrimidine bases and nitroxide spin labels were also explored.^{47,523,524} The first spin-labeled phosphoramidite for incorporation into DNA was the uridine analogue 172, which was prepared by Sonogashira cross-coupling,⁵²⁵ which can be performed during solid-phase synthesis.⁵²⁶ The alkyne linkage limits the flexibility of this probe, making it valuable for DEER applications to measure long-range distances with narrow distance distributions.^{527,528} The cytidine analogue 173 was also described.⁵²⁹ Sonogashira coupling yielding 172 and 173 can also be performed directly during the synthesis of oligonucleotides,^{527,530,531} similar to the attachment of 27 to 2-iodo-adenine (Figure 23).⁵¹⁰ Nucleotide 174^{532,533} is based on a very similar design and proven to be useful in DEER experiments.

As an alternative, the isoindoline based spin labels ^{Im}U, ^{Ox}U, and ^{ExIm}U have been described (Figure 24).^{534,535} Their nitroxide group is aligned with the axis of the rotatable linker to the nucleoside, which has obvious advantages for immobilization and distance measurements by DEER.⁵³⁶ A 2'-methoxy derivative, ^{Im}Um, has also been described and used for distance measurements in RNA duplexes.⁵³⁷ Recently, the tetraethyl nitroxide version ^E^{Im}Um was developed, which is much more resistant to reduction and was, thus, successfully used to study RNA duplexes in oocytes.⁵³⁸

Hopkins and co-workers developed the nucleotide **Q** (Figure 24).^{539,540} Pairing of **Q** with the noncanonical base 2-aminopyridine (2AP) results in a rigid spin label that has been used to study the sequence-dependent dynamics of duplex

DNAs.^{541,542} Sigurdsson and co-workers developed this concept further by designing the rigid spin label **Ç**, which does not require a noncanonical base for pairing, broadening its range of applications (Figure 24).^{543,544} **Ç** represents the fusion of a cytosine with a nitroxide featured isoindole via an oxazine linkage. It is identical with the noncovalent spin label **ç** (Figure 15) except for the linkage to the ribose. **Ç** prefers to pair with guanine and does not perturb the structure of a DNA duplex, as observed in a crystal structure (Figure 25).⁵⁴⁵ A ribo-analogue with a methoxy modification in the 2' position (**Çm**) has been developed for use in RNA.⁵⁴⁶ The synthetic incorporation of **Ç** and **Çm** into DNA and RNA was recently improved to circumvent partial reduction during oligonucleotide synthesis.⁵⁴⁷ Both labels deliver accurate distances in DNA and RNA and have been used in numerous applications.^{548–558} Recently, the tetraethyl nitroxide versions **EÇ** and **EÇm** have been developed and incorporated into DNA and RNA.⁵⁵⁹ They are more inert toward reduction by ascorbate and thus potentially allow in-cell EPR measurements, as recently demonstrated for **E**^{Im}Um.⁵³⁸

Commercially available EDTA-derivatized deoxythymidine (EDTA-C₂-dT) can be incorporated into DNA during oligonucleotide synthesis.⁵⁶⁰ Loaded with a Fe(II) or Mn(II) ion, it has been used to study protein-DNA interactions by NMR measurements of PREs.^{560,561} The long and flexible linker and the occurrence of enantiomeric forms of metal-EDTA complexes challenge applications that require the measurement of PCSs. The design of a wider range of oligonucleotide tags that can be used to measure PCSs is still a little explored area of research.

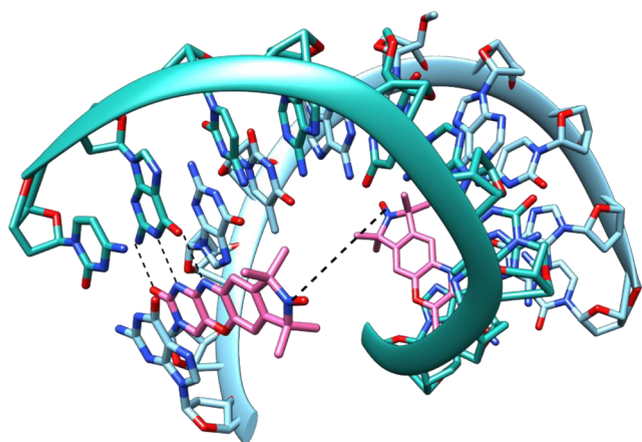


Figure 25. X-ray crystal structure (PDB code 3OT0)⁵⁴⁵ of a 10 base pair DNA duplex containing the ζ spin label (highlighted in pink). Two uridine residues are 2'-methylated. The phosphate backbone is shown as a ribbon. The distance between both nitroxide radicals (18 Å) is indicated. Hydrogen bonds between guanine and ζ are indicated by dashed lines.

Saxena and co-workers incorporated the Cu(II)-binding and commercially available 2,2'-dipicolylamine (dPA) phosphoramidite in lieu of a nucleotide into DNA (Figure 24).^{562–564} Its use in DNA duplexes requires an abasic site in the complementary strand and allows distance measurements by

EPR spectroscopy. A related spin label (L) was used to label DNA quadruplexes with Cu(II) forming a Cu(pyridine)₄ complex (Figure 24).^{565,566} While initial studies used the racemic form of the tag, later studies employed the (S) isomer⁵⁶⁶ for distance measurements. A DNA quadruplex can also be labeled with a large dinuclear platinum(II) complex linked to TEMPO.⁵⁶⁷

3.5. Paramagnetic Probes for Oligosaccharides

Carbohydrates present an additional class of biological macromolecules that have been studied with various paramagnetic tags. To date, the variety of tags and attachment chemistries has been more limited than for proteins and oligonucleotides.^{38,39} Most strategies capitalize on the possibility of selective amination at the reducing terminus using ammonium formate and subsequent formation of an amide bond between tag and oligosaccharide amine (Figure 26). Following this approach, TEMPO has been installed at the reducing end of *N*-acetylglucosamine (Figure 26A, 175) to study its interaction with proteins by NMR spectroscopy.^{568,569} Subsequently, a more complex high-mannose-type oligosaccharide was studied with the same approach.⁵⁷⁰

An EDTA-based lanthanide tag was first attached to the aminated reducing terminus of *N,N'*-diacetylchitobiose (Figure 26B, 176) to enable paramagnetic NMR measurements with an excellent correlation between experimental and predicted PCSs.⁵⁷¹ Subsequently, this tagging strategy was also applied to high-mannose-type oligosaccharides.^{572,573} Biphenyl variants

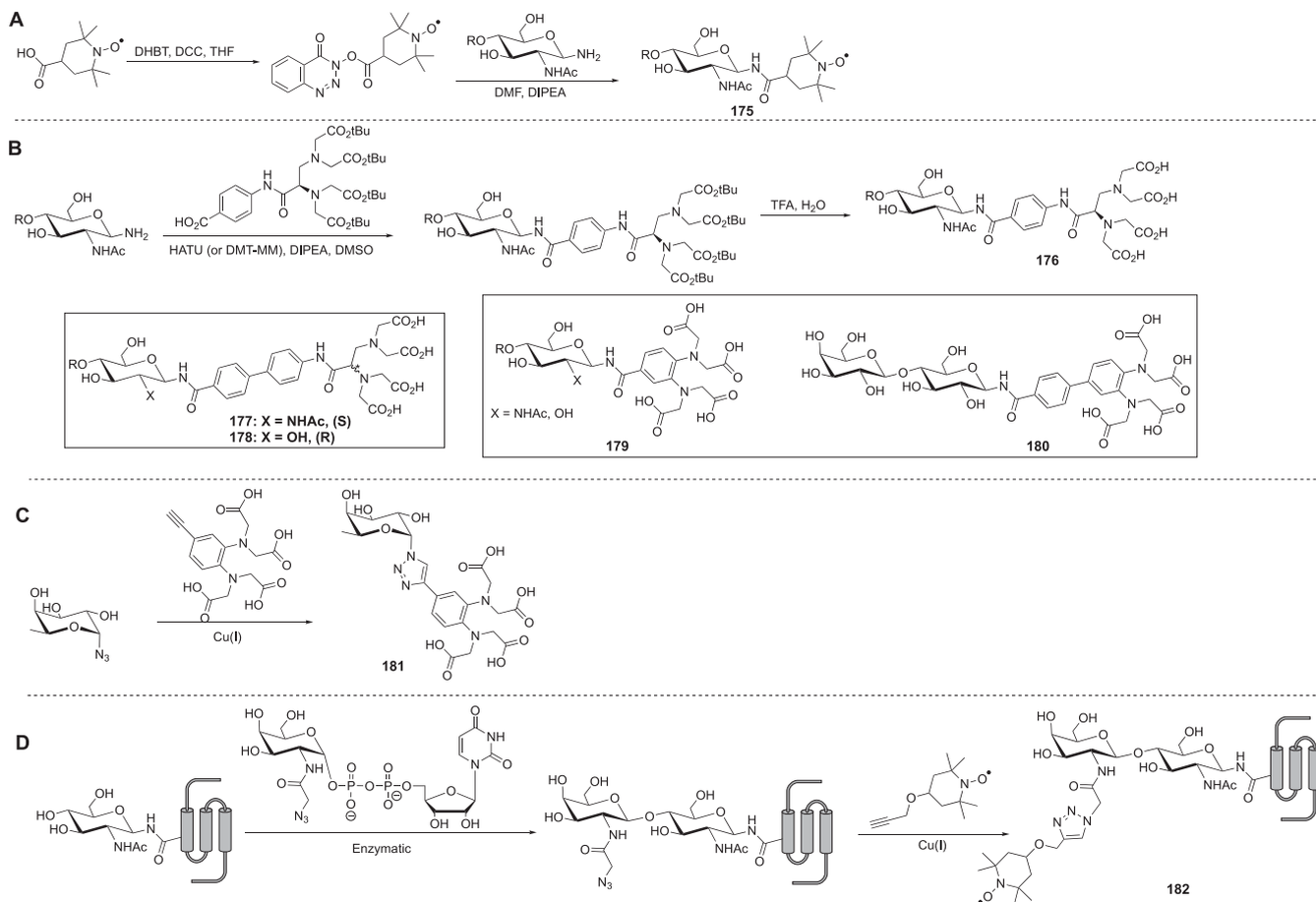


Figure 26. Carbohydrate probes and their synthesis.

of this tag in (S) (Figure 26B, 177)⁵⁷⁴ and (R) (Figure 26B, 178)⁵⁷⁵ configurations were also described.

Kato and co-workers developed a phenylenediamine based lanthanide chelating tag that can be coupled to the aminated reducing terminus of oligosaccharides using standard amide bond formation (i.e., HATU (2-(7-azabenzotriazol-1-yl)-*N,N,N',N'*-tetramethyluronium hexafluorophosphate), DIPEA (*N,N*-diisopropylethylamine), and DMF (*N,N*-dimethylformamide)) yielding **179** (Figure 26B).⁵⁷⁶ This tag has since been used in various applications to study carbohydrates and their interactions with proteins by paramagnetic NMR spectroscopy.^{577–580} A biphenyl variant attached to lactose (Figure 26B, **180**) has also been used to study lactose conformations and its interactions with a receptor protein.⁵⁸¹

Another report described the use of an L-fucose azide and an alkyne based tag for direct “click” reaction to furnish compound **181** (Figure 26C).⁵⁸² It provided an elegant way to obtain a model for the binding site of the fucose on a 70 kDa receptor protein by the use of intermolecular PREs and PCSs to position the lanthanoid ion on the carbohydrate with respect to the protein. In a partially enzymatic and partially synthetic approach, an azide bearing saccharide was selectively introduced to a glycosylation site in a protein and subsequently reacted with an alkyne based label to yield the spin-labeled glycoprotein **182** (Figure 26D).⁵⁸³

4. COMPLICATIONS IN PRE-TO-DISTANCE CONVERSION

The PRE is very strong for a nucleus *I* close to a paramagnetic center *S* and drops off rapidly with increasing distance r_{IS} because of its r_{IS}^{-6} dependence (eqs 9–13). Therefore, PREs are powerful tools for the detection of minor states or the characterization of ensembles of conformations, such as found in intrinsically disordered proteins or encounter complexes.^{33,34,584} At the same time, the strong distance dependence of the PRE adds uncertainties to distance measurements of major conformations, when minor conformations or alternative interactions featuring short distances r_{IS} contribute disproportionately to the observed PRE. Minor species populated as low as 0.1% can be manifested in the observed PRE in this way.

Most obviously, problems arise when proteins are prone to aggregation, but the potential for intermolecular associations may also be introduced by the chemical tag. In principle, intermolecular effects can be detected by measuring the PREs at different concentrations of the tagged molecule,³⁶⁵ but accurate measurements of PREs at low concentration may be very time-consuming for biological macromolecules. Intermolecular PREs can be suppressed by dilution of the paramagnetically labeled molecule with the diamagnetic reference, but as the chemical shifts are conserved in the presence of paramagnetic centers with slow electronic relaxation times, the superimposition of NMR signals from diamagnetic and paramagnetic molecules makes it difficult to disentangle the intra- and intermolecular PREs.

A second complication arises from incomplete paramagnetic tagging. Tags with limited metal binding affinity are easily under- or overtitrated with metal ions and nitroxide tags may be subject to partial reduction or oxidation to diamagnetic species, leading to mixtures of paramagnetic and diamagnetic species even following quantitative chemical ligation. These complications may explain, why the correlation of PREs with distance has been disappointing in many published examples, despite the steep distance dependence of the PREs, compromising the accurate conversion of PREs into distances.^{130,365,434,459,461,585–587}

Finally, the PRE-to-distance conversion is compromised by any flexibility of the paramagnetic tag as well as uncertainties in electronic relaxation rates. In the case of tags containing Gd(III), Mn(II), or Cu(II) ions as the paramagnetic center, the PRE is entirely or predominantly driven by the SBM mechanism, which depends on the effective correlation time of the vector connecting the nuclear spin with the paramagnetic center. In addition, the electronic relaxation rate depends on the ligand-field of the metal complex. In practice, conversion of the PREs into distances, thus, requires calibration against known distances to atoms in the target molecule.^{249,588} In the case of nitroxide tags, the electronic relaxation time is very long (of the order of 100 ns),⁵⁸⁹ but the generally hydrophobic nature of these tags makes the assumption of absence of intra- and intermolecular associations more tenuous.

The problem of incomplete paramagnetic tagging and uncontrolled intermolecular PREs can be solved by using metal tags with anisotropic magnetic susceptibility tensor. In this case, the NMR signals of paramagnetic and diamagnetic species are separated by PCSs. Furthermore, intermolecular PREs can readily be accounted for by mixing the paramagnetically tagged molecule with the diamagnetic reference. In the mixture, the relaxation rates of diamagnetic and paramagnetic species are equally affected by intermolecular PREs, so that the intermolecular effects drop out when calculating the difference in relaxation rates between paramagnetic and diamagnetic species. The concept was explored and demonstrated with calbindin D_{9k} containing Er(III) or other paramagnetic lanthanoid ions as the paramagnetic center and Lu(III) as the diamagnetic reference.¹²⁴ The results showed that PREs extracted from longitudinal relaxation rates were less sensitive to intermolecular effects than transverse PREs but that more accurate distance restraints were obtained by measuring transverse relaxation rates as measurements of R_1 (¹H) relaxation rates are affected by multiexponential relaxation due to cross-relaxation. Specifically, transverse PREs measured with the Er(III) sample delivered reliable distances in the range between 12 and 25 Å with an RMSD below 1 Å, if RDCs in the paramagnetic state were controlled by using short relaxation delays and a magnetic field strength not above 14.1 T. In the case of flexible tags, however, distance variations and difficulties to determine the effective correlation time remain compromising factors in quantitative PRE-to-distance conversions.

5. EXAMPLES OF APPLICATIONS OF PARAMAGNETIC NMR IN PROTEIN STUDIES

Paramagnetic effects provide structural information, which can be used further to assess the dynamics of biological macromolecules, such as domain reorientation or the exchange with minor conformational species. The following section serves to give examples that illustrate the range of applications of paramagnetic probes, including protein resonance assignment, 3D structure determination, studies of protein–protein and protein–ligand complexes, and the identification of lowly populated states. Section 6 adds further examples for an overview of the breadth of applications for the different types of probes.

5.1. Protein Structure Studies

5.1.1. NMR Resonance Assignments in the Paramagnetic and Diamagnetic States. Sequence-specific resonance assignments are the starting point for any protein structural study by NMR spectroscopy. If the protein is not too

large and amenable to uniform labeling with ^{15}N , ^{13}C , and ^2H , the resonance assignment generally is easiest by heteronuclear multidimensional (3D/4D) NMR experiments. For large proteins, complete resonance assignments become challenging because of increasing line width and spectral overlap. PCSs offer a way to increase the spectral dispersion and thus resolve spectral overlap. This is particularly useful for correlation spectra of methyl groups, which otherwise show limited spectral resolution.³⁵⁸

If the 3D structure of the target protein is known in advance, PCSs generated by a site-specifically attached probe can be used to assist the resonance assignments.⁵⁹⁰ Fundamentally, measuring PCSs requires attributing the chemical shift observed for a nucleus in the paramagnetic state to its chemical shift in the diamagnetic state. Different methods have been devised to achieve this.⁵⁹¹ Most commonly, 2D correlation spectra, such as heteronuclear single-quantum correlation (HSQC) spectra, are used, where the cross-peaks of paramagnetic and diamagnetic samples are displaced along approximately parallel lines. This observation holds true in particular for paramagnetic metal ions featuring large $\Delta\chi$ tensors as, in this case, PREs broaden the signals of nuclear spins near the metal ion beyond detection and PCSs can be observed only for nuclei beyond a minimal distance from the paramagnetic center. As the distance between two adjacent nuclear spins is much shorter than the distance from the paramagnetic center, very similar PCSs are observed for both spins involved in a cross-peak in 2D correlation spectra or spectra of higher dimensionality, leading to the cross-peak displacements along approximately parallel lines. This greatly facilitates assigning the paramagnetic peaks to their parents in the spectrum recorded of the diamagnetic reference and the attribution can further be underpinned by using two paramagnetic samples produced with the same probe but different paramagnetic metal ions, which generate PCSs of different magnitude. In this case, all three peaks (two paramagnetic, one diamagnetic) produced between a pair of nuclear spins can be expected to lie on the same line. It is further helpful that, for many probes, the $\Delta\chi$ tensors of Tb(III) and Tm(III) ions tend to shift the cross-peaks in opposite directions relative to the diamagnetic reference (Table 2).^{290,158,258,265} Because the PCSs differ for different protons in a molecule, spectral overlap makes it very difficult to measure proton PCSs by 1D NMR spectroscopy.

In an unbiased way, the assignment of paramagnetic peaks to their diamagnetic parent can be achieved by a spectrum that generates cross-peaks between both. This requires a sample simultaneously containing both the paramagnetic and diamagnetic species, where the probes installed bind the metal ions in a way that allows chemical exchange between paramagnetic and diamagnetic metal ions in the slow-exchange regime (exchange rate between about 1 and 100 s^{-1}). These stringent criteria have been fulfilled in some examples, such as probes **125**³²⁷ and **126**,³²⁸ but are difficult to design in advance.^{330,592,593} Notably, the situation of fast chemical exchange also provides a straightforward way of measuring PCSs as, in this case, they increase continuously with increasing concentration of the paramagnetic metal probe.^{365,591} Lesser binding affinity, however, comes with the risk of lesser binding specificity, which can make it difficult to interpret the PCSs.

Having measured a sufficient number of PCSs to fit a $\Delta\chi$ tensor to the 3D structure of the biological macromolecule, the $\Delta\chi$ tensor can be used to predict further PCSs to assist with the resonance assignment. When using a double-armed probe like

CLaNP-5 (Figure 9, 79) that is relatively rigid relative to the protein, it is also possible to predict an initial $\Delta\chi$ tensor with good accuracy.¹²⁷ As nuclei located at different positions relative to the paramagnetic center can have the same PCS, the distinction needs to be made by additional data, such as PREs, RDCs, and CCR effects,⁵⁷ or by PCSs generated from more paramagnetic centers installed at different sites of the protein. The software packages PLATYPUS,⁵⁷ Echidna,⁵⁹⁴ Possum,⁵⁹⁵ and PARAssign⁵⁹⁶ were developed specifically for resonance assignment using paramagnetic restraints.

An instructive example is the assignment of the diamagnetic resonances of methyl groups of the N-terminal domain of Hsp90 with CLaNP-5 tags (Figure 9, 79) installed at three different sites. The protein contains 76 methyl groups in leucine, isoleucine, and valine residues, and over 60% of them could be assigned based on the protein structure and ^1H PCSs measured in ^{13}C -HSQC spectra with Yb(III) as the paramagnetic metal ion, using PARAssign without the use of any prior assignments from the diamagnetic sample.⁵⁹⁷ For the remaining methyl groups the choice of possible assignments was greatly reduced. As the $\Delta\chi$ tensor orientations of the S50C/D54C and T149C/I187C mutants happened to be almost parallel, the information content was greatly reduced and the assignment was predominantly based on two $\Delta\chi$ tensors only. If the $\Delta\chi$ tensor has been determined independently, for example, from ^{15}N -HSQC spectra of backbone amides, PCSs from a single site with a single paramagnetic metal ion can be sufficient to assign 75% of the methyl resonances in the diamagnetic state using the program Possum, as shown for the 30 kDa complex between the N-terminal exonuclease domain ϵ 186 and the subunit θ of the *E. coli* DNA polymerase III (ϵ 186/ θ).⁵⁹⁵ By starting from a X-ray crystal structure, the work also highlighted the fundamental problem of this approach, which arises when subtle differences exist between crystal and solution structures.

5.1.2. Protein Structure Determination. PCSs arguably deliver the most useful paramagnetic structure restraints for 3D structure determinations of proteins, because they are readily measured with high accuracy. In addition, RDCs obtained by paramagnetic alignment convey useful information about relative domain orientations. Paramagnetically generated molecular alignment and measurement of RDCs also is of interest for studies of protein dynamics,^{598,599} which rely on multiple different alignment tensors, as it can be challenging to obtain a larger number of significantly different alignments with the help of conventional alignment media, which act mostly by steric or electrostatic effects.⁶⁰⁰ Software packages like CYANA,⁶⁰¹ Xplor-NIH,^{70,602} and Rosetta^{603–607} have been extended with modules for protein structure computation with paramagnetic restraints.

In the 1990s, Bertini and co-workers refined 3D structures of proteins with the help of paramagnetic data, demonstrating that the structural restraints gained by paramagnetism can outweigh the loss of NMR signals from ^1H spins in the vicinity of the paramagnetic center.^{112,139} More recently, the benefit of paramagnetic data for 3D structure determinations was highlighted by Nietlispach and co-workers, who applied PCS restraints to refine the structure of the phototactic receptor sensory rhodopsin II (pSRII), a membrane protein with over 240 residues from the family of seven transmembrane helix proteins.⁶⁰⁸ In previous work, the authors had determined the structure of pSRII in a micelle using NOEs.⁶⁰⁹ By labeling pSRII at four different positions with the paramagnetic probe C2 (Figure 9, 96) loaded with different paramagnetic metal ions,

the global fold of the protein could be derived by the exclusive use of PCS restraints. By combining the PCS and NOE data, the structure was refined to a backbone RMSD of 1.5 Å to the mean.⁶⁰⁸

Besides delivering unique long-range structural restraints, the chemical shift measurements needed to measure PCSs can be performed very quickly. This has recently been exploited to elucidate the structure of the transient enzyme intermediate formed by SrtA with its substrate peptide.²⁶⁶ The enzyme–substrate complex is an unstable intermediate that contains a labile thioester bond with the active-site cysteine residue (Cys184) and presents only a minor species in solution. Nonetheless, comparison with the chemical shifts of a stable disulfide-bond-linked analogue enabled the resonance assignment of the thioester intermediate and attachment of the probes 76 and 77 (Figure 7) loaded with one of four lanthanoids, Dy(III), Tb(III), Tm(III), and Lu(III) to SrtA delivered PCSs. Using the PCSs as input for Xplor-NIH, the structure of the thioester intermediate was calculated as an ensemble of conformers with a RMSD of 1.1 Å.²⁶⁶

The sensitivity with which PCSs can be measured is also of great advantage for measurements at low concentrations, such as encountered in in-cell studies of protein structure. Using GB1 as a model protein, Müntener et al. attached the M7Py-DOTA tag (Figure 9, 88) loaded with one of three lanthanoids, Tb(III), Tm(III), and Lu(III), and recorded PCSs and RDCs at about 50 μM protein concentration after transferring the labeled protein into *Xenopus laevis* oocytes.²⁸⁶ The paramagnetic restraints measured by in-cell NMR assisted the 3D structure determination using GPS-Rosetta.⁶⁰⁶ The same protein at the same low concentration was targeted at the same time by Pan et al., using PCSs measured in *X. laevis* oocytes.⁶¹⁰ To measure PCSs, this group attached 4PS-PyMTA (Figure 7, 76) loaded with Tb(III), Tm(III), Yb(III), or Y(III) ions and calculated the structure without RDCs using GPS-Rosetta. The authors were careful to point out, however, that Rosetta⁶¹¹ is one of the most powerful software packages for 3D structure predictions of small proteins from the amino acid sequence alone.⁶¹⁰ Specifically, they showed that the fold of GB1 is already correctly predicted by CS-Rosetta,⁶⁰³ when backbone chemical shifts are provided as the only experimental data.⁶¹⁰

The conversion of PCS data into structure restraints requires knowledge of the $\Delta\chi$ tensor, which is best obtained by fitting to 3D structure coordinates. For proteins of unknown structure, the powerful model building algorithm provided by Rosetta thus is an excellent starting point where models that do not allow good $\Delta\chi$ tensor fits can be discarded early in the structure calculation.⁶⁰³ In this way, PCS data not only improve the quality of the final structures but also accelerate the convergence of the calculations. The value of PCS restraints for protein fold determination, even if backbone chemical shifts and PCSs from backbone amide protons are the only experimental data provided, is very clear for larger proteins.^{604,605,607,612}

Because of their long-range nature, PCSs are particularly suited to determine full structures in situations, where the structure of a rigid protein domain is known while the conformation or orientation of another part is not. In this situation, even sparse PCS data have been shown to be sufficient to fit $\Delta\chi$ tensors to the rigid part and use the PCSs of the unknown part to determine its structure or relative orientation.^{147,291,612–619}

It is instructive to compare the PCS structure restraints with the way, in which the global positioning system (GPS) identifies

the location of objects on Earth. Just as the measurement of the distances to several satellites identifies a single location on the surface of Earth, the PCSs of a nuclear spin can be measured for different protein samples prepared with tags positioned at different sites. Each PCS defines an isosurface and the intersection of four PCS isosurfaces defines a single point in space.^{606,620} The concept was demonstrated with the 3D structure determination of the C-terminal domain of ERp29 tagged at four different sites either with IDA-SH (Figure 11, 126) or C1 tags (Figure 9, 96), using Tm(III) and Tb(III) as the paramagnetic ions.⁶⁰⁶ The same concept was applied to determine the structure of this protein with the help of PCSs measured with Co(II) ions bound to double-histidine motives in four different helices, one at a time.⁵⁹² Notably, PCSs of the backbone amide protons alone were sufficient to obtain the correct protein fold. While PCSs from an increasing number of tagging sites increases the level of structural information available, structural details on the conformations of amino acid side chains can already be obtained with two different tagging sites, as the side chain locations are restricted by their link to the backbone.^{94,621}

5.1.3. Protein–Protein Interactions. With a paramagnetic probe on one of the interacting proteins in a protein–protein complex, the paramagnetic effects in the partner protein can be recorded to determine the relative position and orientation of the two proteins in a rigid body docking approach, which assumes that the structures of the individual binding partners are conserved in the complex (Figure 27). The first example of this approach, from 1998, used the paramagnetic heme in cytochrome *f* to induce PCS in the electron transfer partner plastocyanin and model the structure of the complex.^{622–624} In an early example using lanthanoid ions, the structure of the ϵ 186/ θ complex was modeled with the help of PCSs measured with Dy(III), Ho(III), or Er(III) data in the natural metal binding site of ϵ 186, using a rigid body docking approach, although the PCSs were also used to refine the structure of θ in its bound state.^{69,625}

PCSs were also successfully used to model the structure of a homodimer. In this case, two mutants of p62 PB1, named DR and KE, were produced to prevent oligomerization beyond dimer formation.¹⁵⁹ A two-point anchored LBP was incorporated at the N-terminus of DR to coordinate with paramagnetic lanthanoid ions. The lanthanoid position relative to the DR mutant was determined by fitting the $\Delta\chi$ tensors and PCSs were measured in the KE mutant. Using the measured PCS and chemical shift changes observed at the interface of the KE mutant, rigid body minimization protocol in Xplor-NIH^{70,602} delivered the structure of the dimer. It was noted, however, that multiple solutions were possible even with the availability of PCS data sets from different lanthanoid ions as the $\Delta\chi$ tensor axes tended to be similar. The degeneracy could be resolved by modifying the LBP.⁶²⁶

Protein–protein interactions (PPI) are based on noncovalent forces, such as charge–charge attractions, hydrophobic interactions, and hydrogen bonds, and paramagnetic effects are excellent tools to investigate very weak protein–protein interactions such as observed in encounter complexes. As the protein partners approach each other in solution, the formation of an encounter complex precedes the formation of the specific protein–protein complex. These weak and dynamic complexes are essential for rapid complex formation, yet difficult to characterize by traditional techniques.^{627,628} PREs are particularly suited for investigating such weak PPIs, as they are

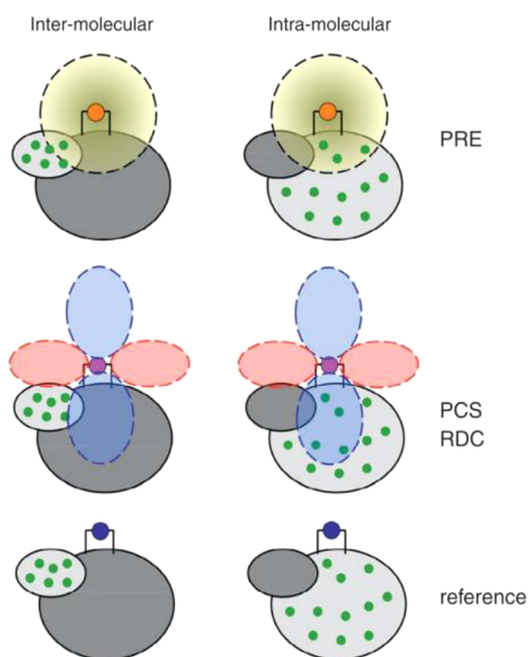


Figure 27. Strategy for using PREs, PCSs, and RDCs for structure determination of protein–protein complexes using differently labeled samples. PREs are measured using a tag with slow electronic relaxation (e.g., Gd(III)) or a paramagnetic center with little χ tensor anisotropy (e.g., Er(III)). PCSs and RDCs are obtained using the tag with an anisotropic paramagnetic center (e.g., Tm(III)). The data are measured with respect to a diamagnetic reference produced with the tag containing, for example, a Lu(III) ion. To simplify the NMR spectra and distinguish between intermolecular and intramolecular effects, the complex can be prepared with isotope labeling (symbolized by green dots) of one of the proteins only. The intramolecular restraints are used to fit the position of the paramagnetic center and orient the $\Delta\chi$ tensor. 2D NMR spectra (typically $[^{15}\text{N},^1\text{H}]$ -HSQC spectra) can be sufficient to obtain a complete set of inter- and intramolecular PREs, RDCs, and PCSs, which allow determining the relative position and orientation of the two proteins in a rigid body docking approach. Reproduced with permission from ref 30. Copyright 2014 Elsevier.

insensitive to the relative orientation of the proteins and sensitive to minor states in which a nucleus is closer to a paramagnetic center than in the major state, which usually is the final specific complex. Complementary information can be obtained by PCSs and RDCs, provided that the orientation of the $\Delta\chi$ tensor changes only over a limited angular range, when the proteins populate an ensemble of complex structures. Averaging over a greater range is evidenced by averaging of the PCSs and RDCs to zero.

The electron-transfer complex of yeast iso-1-cytochrome *c* (Cc) and yeast cytochrome *c* peroxidase (CcP) is an example of a weak PPI comprehensively characterized by paramagnetic NMR. Both proteins contain a paramagnetic heme group, but the intermolecular PCSs are too small to be suitable for structural studies. Following the attachment of a paramagnetic tag (MTSL, Figure 3) at different locations on the CcP surface, PREs were recorded for Cc and utilized to dock Cc on CcP. The resulting model of the complex was similar to the crystal structure. In this case, the encounter complex represented no less than 30% of the complex and its interface was preferentially located near that of the final specific complex.^{65,91} CcP variants with added negative patches were used to study the role of

charge distribution and PREs showed that cytochrome *c* visits such new patches in the encounter complex.⁸³

In another example, the complex between cytochrome P450cam (cytP450cam) and putidaredoxin (Pdx) was investigated using CLaNP-7 (Figure 9, 80) loaded with Tm(III) or Gd(III) to obtain intermolecular PCSs, RDCs, and PREs.⁶²⁹ The position of putidaredoxin on cytP450cam found with these restraints was in good agreement with crystal structures obtained independently.^{629,630} The study highlighted the value in combining different types of paramagnetic information to obtain a good structure. In addition, the NMR data provided evidence for the presence of a lowly populated encounter complex. Using additional paramagnetic restraints and the maximum occurrence of regions methodology,⁶³¹ it could be shown that in this encounter complex Pdx visits a large area of the surface of cytP450cam.

With increasing molecular weight of the system, it becomes increasingly difficult to obtain the specific NMR resonance assignments required to fit $\Delta\chi$ tensors. In this situation, site-specific incorporation of a probe that can easily be observed in the NMR spectrum can be combined with a paramagnetic probe to monitor the PRE on the NMR probe and, thus, derive distance restraints. The concept has recently been demonstrated by Abdelkader et al., who used genetic encoding for site-specific incorporation of an unnatural amino acid carrying a trimethylsilyl (TMS) group at the end of the amino acid side chain to obtain a readily observable ^1H NMR probe (an intense singlet near 0 ppm).⁶³² Using a 71 kDa homodimeric *p*-cyano-*L*-phenylalanyl-tRNA synthetase as an example, a mixture of TMS labeled protein and paramagnetically labeled protein (using either MTSL or the C1 tag loaded with Gd(III) attached to a cysteine residue) revealed a PRE across the dimer interface with Gd(III) but not with MTSL.⁶³² Taking into account the flexibilities of the tags and TMS amino acid side chain, the data could be interpreted as a distance restraint between about 10 and 20 Å.

5.1.4. Protein–Ligand Interactions. NMR spectroscopy is uniquely suited to study protein–ligand interactions in drug discovery campaigns because NMR works in solution and can provide atomic level information about the binding site and ligand orientation. It is useful to distinguish between the scenarios where the exchange rate of the ligand is fast or slow on the time scale of the NMR experiment. Rapid and slow ligand binding and dissociation generally correlates with relatively weak and strong ligand affinities, respectively. Paramagnetic NMR opens new approaches for studies in either limit.

Fragment-based ligand screening is a method that searches for small molecules that bind to proteins weakly. The use of paramagnetic NMR for fragment-based ligand screening has been explored by Saio et al. with a double-anchored LBP.⁶³³ Binding of the ligand was reported by PREs generated by a Gd(III) ion in the LBP, where the PREs were detected as increased transverse relaxation rates of the ^1H signals in 1D NMR spectra of the ligand, using $T_{1\rho}$ relaxation experiments. In the limit where the ligand exchanges between free and bound state faster than the transverse relaxation rate, the relaxation enhancement experienced in the bound state due to the proximity of the paramagnetic center in the target protein is transferred to the spectrum of the free ligand. In analogy to the transferred NOEs and saturation transfer, the effect is referred to as transferred PRE (tPRE). The size of a tPRE reflects the distance between the nuclear spin of the ligand in the bound state and the paramagnetic center attached to the target protein.

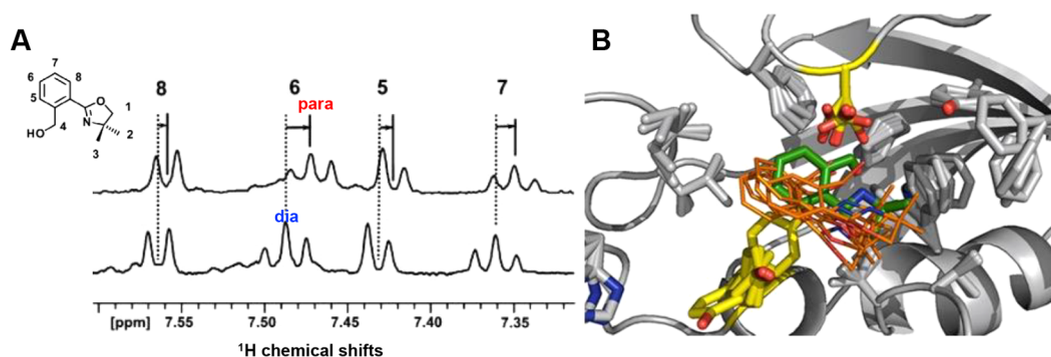


Figure 28. Transferred PCSs identify the binding site and binding orientation of ligand molecules that are in fast exchange with excess free ligand. (A) tPCSs (the difference between the resonance positions for the paramagnetic (para) and diamagnetic (dia) samples) observed in the ^1H 1D NMR spectra of excess ligand in complex with the protein, which is tagged with either paramagnetic or diamagnetic ions (one site at a time). The dashed and solid lines identify the ligand resonances in the diamagnetic and paramagnetic samples, respectively. (B) Superposition of the averaged NOE structure (in green) of the FKBP12–1 complex and the best five structures (in orange) calculated using tPCSs. The protein backbone is represented as a gray ribbon except for the residues Asp37 and Tyr82 (yellow). The average RMSD of the ligand from PCS calculations relative to the NOE calculation is 2.8 ± 0.4 Å. Reproduced with permission from ref 620. Copyright 2013. American Chemical Society.

In the same way, also transferred PCSs (tPCS) can be observed and used to assess the structure of protein–ligand complexes.⁶³³

In another example, Guan et al. demonstrated that the location and orientation of a ligand can be determined solely using tPCSs, provided that the ligand is in fast exchange between bound and free states relative to the size of the PCS (expressed in rad s^{-1}).⁶²⁰ In the fast exchange limit, the experiments allow using the free ligand in excess, that is, only a small fraction of the ligand is bound and the PCSs measured are scaled by the bound fraction. Although the PCSs measured are thus relatively small, the smaller size is more than compensated by the narrow line width observed for the ligand, which is dominated by the fraction of free ligand. Placing the paramagnetic center, Ln(III)-CLaNP-5 (Figure 9, 79), at three positions, one at a time, on the target protein FKBP12 (Figure 28) enabled a PCS-based triangulation approach to capture the ligand's site of binding and orientation relative to the protein. The structure of the complex obtained from tPCSs agreed with the restraints obtained from NOEs but was easier to obtain in the absence of extensive ^1H NMR resonance assignments of the protein.

In an early example, John et al. showed that tPCSs generated by a single lanthanide binding site can be sufficient to determine the binding site and orientation of a weakly bound ligand, if steric restrictions imposed by the protein invalidate any of the alternative structural solutions of the tPCS data set.⁶³⁴ In this example, the lanthanide ion was bound deep inside the metal binding pocket of the $\epsilon 186/\theta$ complex. The ligand (thymidine) bound with millimolar binding affinity and tPCSs were observable in 1D ^1H and ^{13}C NMR spectra of the ligand present in over 500-fold excess over the protein complex. The sensitivity of the approach is illustrated by the speed (<0.5 h) with which the ^{13}C NMR spectra were recorded at natural isotopic abundance. The binding mode coincided closely with that observed in the crystal structure of the $\epsilon 186$ –thymidine monophosphate complex.

A caveat associated with tPRE and tPCS effects is the possibility that the weakly binding ligand temporarily associates with the paramagnetic tag rather than the target molecule. False positives arising in this way cannot adequately be addressed by a control experiment probing the direct interaction between ligand and free paramagnetic tag, as unexpected interactions are likely promoted by the formation of binding pockets between

tag and target molecule. The strong distance dependence of PCSs and PREs notably amplifies the effects of minor populations of direct ligand–tag interactions.

As ligand binding tends to affect the global structure of the target molecule little or not at all, changes in PCSs observed locally in the target molecule can be used to determine local structural changes arising from the interaction with a ligand, such as changes in amino acid side chains. A recent study labeled the protein FKBP12 with Ln(III)-CLaNP-5 (Figure 9, 79) at three sites, and valine and leucine residues were selectively isotope labeled with $^{13}\text{CH}_3$ groups.⁶²¹ Significant changes in PCSs were observed between the free and ligand-bound protein for methyl groups located near the ligand binding site, indicating displacements of the methyl groups in the 1–5 Å range.

In the slow exchange limit, it is not meaningful to use ligand in excess over the target protein, which makes it much harder to discriminate the NMR signals of the ligand against the background of protein signals, especially as the chemical shifts of the bound ligand usually differ from those in the free state. While isotope labeling of the ligand enables its selective detection by NMR spectroscopy, the chemical synthesis of isotope labeled ligands is cumbersome in practice. Chen et al. demonstrated that chemical labeling of the ligand with a *tert*-butyl group presents a synthetically more practical alternative.⁶³⁵ As the ^1H NMR signal of the *tert*-butyl group is an intense signal, it can readily be detected by 1D ^1H NMR. The specific example used was the 27 kDa complex between the dengue virus NS2B-NS3 protease, tagged at three different sites with C2 tags loaded with Tm(III), Tb(III), or Y(III), and a covalently binding peptide inhibitor labeled with a *tert*-butyl group.⁶³⁵ NOESY spectra revealed PCSs not only of the *tert*-butyl group but also of ligand resonances of protons near the *tert*-butyl group, which enabled positioning and orienting the ligand in the bound state. Although the structural information obtained for the ligand in this way appears limited, it proved decisive in identifying the correct pose obtained by independent modeling of the protein–ligand complex.

5.1.5. Structures of Minor States. Protein folding, enzyme catalysis, and other functions require conformational exchange, involving excited states with low populations and short lifetimes (microseconds to milliseconds). The structures of these lowly populated states can usually not be solved by X-ray

Table 4. Applications of Synthetic Probes in Protein NMR Studies

probe	applications in protein NMR studies	comments
MTSL (Figure 3)	protein global folding and structure determination ^{305,380,699,934,656,666,669,671} protein–protein interaction ^{65,83,91,658} protein–ligand interaction ^{661–663} protein dynamic studies ^{664–667,670,730} protein distance measurements by ¹⁹ F-PRE ^{5,87} protein conformation studies ⁶⁶⁷	small tag generating sizable PRE effects; easy to handle and commercially available; in NMR experiments, this tag produces PREs as the only paramagnetic effect; the nitroxide group is sensitive to reduction or oxidation; reduction of the disulfide bond formed with the target leads to loss of the tag
maleimide-TEMPO (Figure 5, 15)	protein structure determination ^{249,250,643,644,674,675} protein binding target determination ⁶⁷⁶ protein alignment ²⁵¹	commercially available and attached to protein via stable thioether linker
SPy-EDTA and analogues (Figure 7, 60–61)	protein structure determination ²⁶⁶ and in-cell protein structure determination ⁶¹⁰ protein–peptide complex ²⁶³ protein–ligand interactions ^{678,679} protein complex NMR signal assignment ⁶⁷⁷	commercially available, but exists in multiple conformations lanthanoid based probe for attachment to single cysteine residues; small, hydrophilic, and carrying a low net charge; forming a stable thioether linkage with cysteine; complete ligation yields can be difficult to obtain
4MMPyMTA and analogues (Figure 7, 73–77)	methyl group assignment ^{597,621} protein–ligand interaction studies ^{768,820,617,641} protein structure determination ^{616,682,731} protein dynamic studies ^{274,275} protein–protein complex studies ⁶⁸⁰ protein–peptide complex studies ⁶⁸¹ protein complex studies ^{629,631,685–687} protein structure determination ⁶²⁹ protein conformation studies ⁶⁸⁵	frequently used lanthanoid-based cyclen probe, suitable for PCS measurements; the double-armed linkage to the target results in high rigidity; it carries a net charge of 3+ and some of its pendants are fairly hydrophobic; the disulfide linkage between probe and target is prone to reduction
CLaNP-5 (Figure 9, 79)	protein structure determination ^{616,682,731} protein dynamic studies ^{274,275} protein–protein complex studies ⁶⁸⁰ protein–peptide complex studies ⁶⁸¹ protein complex studies ^{629,631,685–687} protein structure determination ⁶²⁹ protein conformation studies ⁶⁸⁵	similar to CLaNP-5, but less charged and with yellow color
DOTA-M8 (Figure 9, 86)	intrinsically disordered proteins; increasing NMR spectral dispersion ⁶⁹⁴ protein domain studies ^{695,696} protein alignment ⁶⁹⁷ protein–ligand interaction ⁶⁹⁸ RNA structure analysis ⁶⁹⁹ protein structure refinement ⁷⁰¹	binds Ln(III) tightly; produces large $\Delta\chi$ tensors, and has zero net charge; disulfide bond after attachment to the protein may be prone to reduction; two conformations were reported.
M8-CAM-1 (Figure 9, 87)	in-cell protein structure determination ²⁸⁵	shares the core structure with DOTA-M8 but a propylidooacetamide attachment replaces the pyridylthio linker to form a more stable C–S linkage; a long tether to the protein backbone reduces its $\Delta\chi$ tensor magnitudes
M7Py-DOXA (Figure 9, 88)	in-cell protein structures from 2D NMR experiments ²⁸⁶ homotrimeric protein structure investigation ⁷⁰⁰ protein–ligand interaction ⁶⁹⁸ protein structure refinement ⁷⁰¹	an analogue of DOTA-M8, with a 4-(phenylsulfonyl)pyridine moiety yielding attachment with a short and rigid linker but the attachment reactivity is low
M7PyThiazol-DOXA (Figure 9, 92)	protein structure determination ⁷⁰¹	a newer version of M7Py-DOXA; reacts with cysteine with higher selectivity and efficiency; generates a $\Delta\chi$ tensor even larger than DOTA-M8
C1/C2 (Figure 9, 96)	methyl group assignment ⁶⁸⁹ protein structure determination ^{94,604,606,612}	lanthanoid based cyclen probe for attachment to single cysteine residues; suitable for PCS measurements but with high net charge and bulky hydrophobic pendants; forming a disulfide linkage between probe and target; C1 and C2 are enantiomers

Table 4. continued

probe	applications in protein NMR studies	comments
C3 (Figure 9, 98)	protein–protein interactions and protein–ligand interactions ^{2,9,10,16,19,690} DNA–protein domain binding studies ⁶⁹³ protein structure studies ⁷⁰²	similar to the C1 and C2 tags but for attachment to <i>p</i> -azido- <i>L</i> -phenylalanine by Cu(I)-catalyzed click chemistry, which is orthogonal to cysteine chemistry commercially available and frequently used but somewhat hydrophobic; unstable under reducing conditions
TEMPOL (Figure 12)	protein accessible surface characterizations ^{345,346,355,711–714,732,733} BPTI aggregation ⁷¹⁵ protein folding ^{3,47} protein dynamics studies ^{664–667,670,730} protein distance measurements by ⁹ F-PRE ³⁸⁷	high solubility, stability, and low protein binding affinity; zero net charge
Gd(III)DTPA-BMA (Figure 13)	protein surface accessible regions characterizations ^{330,716,734} BPTI aggregation ⁷¹⁵ protein structure determination ^{352,721} protein complexes studies ^{719,723} protein domain studies ⁷²² spectral editing ⁷²⁶ protein dynamic determinations ³⁵³	zero net charge
Gd(III)TTHA-TMA (Figure 13)	protein–protein interaction ⁷¹⁸	high negative charge; can generate both PCs (loaded with Tb(III)) and PREs (loaded with Gd(III)) in some cases
[Ln(III)(DOTP)] ⁵⁻ (Figure 13)	multisite protein–ligand binding interactions ²⁴	can generate both PCs (loaded with Tb(III)) and PREs (loaded with Gd(III)) in some cases
[Ln(III)(DPA) ₃] ³⁻ (Figure 13)	cationic peptides structure determination ²⁵ spectral editing ³⁵⁸	negatively charged
[Ln(III)(DTPA)] ²⁻ (Figure 13)	identification of residues close to carboxylates ³⁶⁶	commercially available and zero net charged
Gd(III)(DO3A) (Figure 13)	protein–protein interface mapping ³⁵⁷	commercially available and negatively charged
[Ln(III)(EDTA)] ¹⁻ (Figure 13)	spectral editing ³⁵⁸	

crystallography. Relaxation dispersion measurements using the Carr–Purcell–Meiboom–Gill (CPMG) sequence and $T_{1\rho}$ experiments, as well as chemical exchange saturation transfer (CEST) experiments are NMR methods, which have proven uniquely suitable for the study of such high-energy states of proteins.^{636–638}

Relaxation dispersion experiments can yield the absolute difference in chemical shifts ($\Delta\delta$) of nuclear spins between the minor and major states. In combination with RDC restraints obtained in an external alignment medium, the structure of a minor state can be determined in favorable cases.^{639,640} It is still difficult, however, to convert the limited experimental information into a 3D structure. In this situation, PCSs of the minor state would provide powerful structural restraints. If the PCSs of the minor state differ from those of the major state, they can be extracted from a relaxation dispersion experiment, where the diamagnetic control sample delivers the chemical shift change in the diamagnetic state and the paramagnetic sample delivers the $\Delta\delta$ values including the PCSs.²⁷⁵ For this application, rigidity of the paramagnetic center is critically important to obtain accurate structural restraints. Earlier relaxation dispersion studies of cardiac troponin C substituted with lanthanoids (Ce(III) or Pr(III)) indicated that the PCSs observed most likely reflect motions of the paramagnetic center.⁶⁴¹ To avoid this problem, both the attachment site and the probe must be rigid on the millisecond time scale. When CLaNP-5 loaded with Tm(III), Yb(III), or Lu(III) was linked to Paz and Cc, which are known to be rigid proteins, ^1H CPMG relaxation dispersion experiments revealed dispersion effects likely reflecting movements of the tag rather than protein.²⁷⁵ When CLaNP-5 was attached to adenylate kinase, relaxation dispersion experiments similarly indicated the presence of a minor conformational state of the probe, in which the $\Delta\chi$ tensor is rotated by 20° relative to the major state. This work illustrated the difficulty of finding a probe without any exchange phenomena on the millisecond time scale. In particular, ring flips of the cyclen ring are hard to suppress.²⁷⁴

Alternatively, PCS restraints for determining the structure of a minor state can be derived from CEST experiments. For a proof of concept of PCS-CEST experiments, Ma et al. used Abp1p SH3–Ark1p, which is a protein–peptide complex with a slow off-rate of the bound peptide, as a model system.²⁶³ 4MMPyMTA (Figure 7, 73) loaded with Tb(III) was attached to the SH3 domain and the minor state was produced by adding only 0.03 equiv of Ark1p peptide. The ^{15}N PCSs of the minor species determined by the CEST measurements proved to be in agreement with the 1:1 Abp1p SH3–Ark1p complex. The same paramagnetic tag was attached to the FF domain of HYPB/FBP11, which is in equilibrium with <2% of unfolded conformation. The PCS-CEST experiment reproduced these results. In particular, the experiments showed that the PCSs of the minor species were small, as expected for a less-ordered conformation.

5.2. Biomolecules with Paramagnetic Tags in the Solid State

The protein GB1 has been investigated extensively as a model protein to test the effect of cysteine ligated paramagnetic tags by MAS solid-state NMR.⁶⁴² Longitudinal R_1 PREs of ^{15}N spins in GB1 were measured in the presence of SPy-EDTA (Figure 7, 60)²⁴⁸ loaded with Cu^{2+} ,⁶⁴³ and also with Mn^{2+} .⁶⁴⁴ The PRE data proved to be good indicators of distance and were measured for 6 tagging sites and used as structural restraints for

computationally solving the protein structure,⁶⁴⁵ while also accounting for tag flexibility.⁶⁴⁶ A less flexible short-armed cyclen based tag TETAC (1-(2-(pyridin-2-ylidysulfanyl)ethyl)-1,4,7,10-tetraazacyclododecane) (Figure 9, 112) was also investigated with Cu^{2+} in GB1.³⁸⁸ While the relatively short T_{1e} time of Cu^{2+} ions results in appreciable R_1 PREs by the SBM mechanism, R_2 PREs are often small, which presents the basis for increasing the sensitivity of NMR experiments by accelerating the recovery of longitudinal magnetization between scans with little increase in line width.⁶⁴⁷ R_1 PREs of ^1H spins are challenging to measure in solids due to spin diffusion effects, but R_2 PREs of ^1H can offer long-range distance restraints.⁶⁴⁸ R_2 PREs by the SBM mechanism are largest for paramagnetic centers with slowly relaxing electrons such as the nitroxide spin label, which has also been demonstrated in GB1 by ligating cysteine residues with MTSL.⁶⁴⁹ Measurement of PCSs generated by paramagnetic metal tags is more difficult in solids than in solution, because different orientations of the metal complex relative to the protein would lead to a range of different PCSs. Nonetheless, PCSs have successfully been measured in GB1 ligated with 4MMDPA tag (Figure 11, 116) loaded with Co^{2+} and lanthanoids and were used to help solve the protein structure with Rosetta.³⁸⁹

Because of the close proximity of molecules in the solid state, intermolecular paramagnetic effects can be substantial. To minimize these, in most investigations of paramagnetic GB1 in the solid state the sample was diluted with diamagnetic protein. Alternatively, intermolecular paramagnetic effects in a purely paramagnetic sample can be accounted for by methods of NMR crystallography that consider paramagnetic effects in a periodic lattice.⁶⁵⁰

Aside from solid proteins in the microcrystalline state, paramagnetic effects have also been investigated by solid-state NMR of membrane proteins. For example, the seven-helix transmembrane protein sensory rhodopsin from *Anabaena* tagged at two sites with MTSL nitroxide spin labels was investigated in lipid bilayers.^{651,652} Intermolecular PREs arising from the nitroxide spin label revealed the oligomerization interface of the monomer. Another membrane system, lactose transport protein from *Streptococcus thermophilus* tagged with 3-maleimido-proxyl (Figures 5 and 14),²¹¹ was investigated in native membranes.⁶⁵³ Here, PREs were measured for a ^{13}C -labeled substrate to probe the proximity of an interhelix loop to the protein active site. Finally, PREs have also been utilized to solve the structure of a precipitated protein–RNA complex, named L7Ae, by using IA-PROXYL spin label (Figures 5 and 17)²¹³ at various tagging sites of the protein and measuring intermolecular PREs in isotope labeled RNA.⁶⁵⁴

6. APPLICATIONS OF DIFFERENT TYPES OF PARAMAGNETIC PROBES

Table 4 presents an overview of synthetic probes that have been successfully applied to protein studies by NMR spectroscopy. It provides an impression of the range of applications, where the various probes have been found to be useful. The table also includes some of the probes developed with the aim to obtain narrow distance distributions in DEER experiments. This section briefly summarizes the applications of the different probes listed. Examples that were already mentioned in section 5 are not discussed in detail again here.

6.1. Applications of Nitroxide Probes

Commercially available **MTSL** (Figure 3) has been used successfully in a wide range of protein NMR studies, making it impossible to compile all usage in a comprehensive way. Methods have been devised to speed up accurate ^1H PRE measurements⁶⁵⁵ and been applied to demonstrate that **MTSL** derivatized with a bulky group sufficiently limits the internal motions of the tag to enable the PRE data analysis with the assumption of a single nitroxide position relative to the protein.²¹⁹ The potential of PREs generated by **MTSL** for global fold determination of proteins was assessed by Battiste and Wagner, who showed their value for determining the fold of eukaryotic translation initiation factor 4E (eIF4E) in combination with $\text{H}^{\text{N}}-\text{H}^{\text{N}}$ NOEs and chemical shift data.¹³⁰ Choosing *Bacillus amyloliquefaciens* barnase as the model protein, Gaponenko et al. also provided an early demonstration of the value of **MTSL**-generated PREs in improving the accuracy and ease of global folding determination.⁶⁵⁶ The long-range nature of PREs (generated by **MTSL**) was shown to be particularly valuable for determining the structure of a loop-rich protein, SAPK-interacting protein 1 (Sin1) in combination with limited NOE data.⁶⁵⁷

Examples for the use of **MTSL** in NMR studies of protein–protein complexes have been mentioned in section 5.^{65,91,658} In a recent example, Olivieri et al. used a mixture of ^{15}N - and **MTSL**-labeled ubiquitin with $^{15}\text{N}/^{13}\text{C}$ -labeled ubiquitin to distinguish intra- and intermolecular PREs in the transient diubiquitin complex in a single sample.⁶⁵⁹ **MTSL** has also been used more generally in NMR studies of complexes between proteins and smaller ligands. Gochin and co-workers employed paramagnetic NMR in a second-site screening campaign⁶⁶⁰ to identify ligands binding in the hydrophobic pocket of HIV-1 gp41.⁶⁶¹ By labeling a short peptide, which was known to bind at the edge of the pocket, with **MTSL** or cysteamine-EDTA loaded with either Fe(II) or Co(II), the authors measured the paramagnetic effects on the low-molecular weight ligand as it was bound to the protein, thus detecting its binding site.⁶⁶¹ The **MTSL** labeled peptide alone provided sufficient information as the same group, in a subsequent study, used only **MTSL** labeled peptide to determine the binding site of a different indole ligand binding to HIV-1 gp41.⁶⁶² PREs measured with Rho guanine-nucleotide dissociation inhibitor 2 (RhoGDI2) labeled with **MTSL** provided Ruan and co-workers with key structural information to characterize the binding mode of three hits identified from a fragment library of 890 compounds.⁶⁶³

The strong distance dependence of PREs generated by **MTSL** has been successfully exploited in studies of protein dynamics, including studies of the interdomain movements of cardiac troponin C⁶⁶⁴ and calmodulin.⁶⁶⁵ Using a mixture of **MTSL** labeled and isotope labeled protein, Yang et al. monitored the slow exchange of monomeric units in the homodimer Dsy0195 from *Desulfitobacterium hafniense* Y51 by paramagnetic NMR and DEER experiments.⁶⁶⁶ Investigation of the open-to-closed transition of maltose-binding protein with PREs generated by the probe **maleimide-TEMPO** (Figure 5, 15) revealed that the apo state of the protein populates a major open form and a minor closed species.⁶⁶⁷

MTSL has also been utilized in NMR studies of proteins in the solid state,^{649,652,668,669} where structural restraints from PREs can amount to a sizable fraction of the data available for structure calculations. Especially in the case of membrane proteins, **MTSL** has proven useful to assess their dynamics,⁶⁷⁰ structure,^{586,671} and function.⁶⁷²

Using the **MTSL** tagged and cyanobacterial lectin CV–N-containing fluorinated amino acids as an example, Matei and Gronenborn demonstrated that the high resolution of ^{19}F NMR spectra allows facile measurement of the distances between ^{19}F spins and the paramagnetic center by ^{19}F PREs, which proved to be in a good agreement with the results from ^1H PRE measurements.⁵⁸⁷ Following labeling of a Myc peptide with a CF_3 compound, ^{19}F PREs were also shown to provide a convenient way of assessing proximity to **MTSL** labeled Max peptide, both in the disordered state and the structurally defined complex with DNA.⁶⁷³

6.2. Applications of Aminopoly(Carboxylic Acid)-Based Probes

Dvoretzky et al.⁶⁷⁴ and Vadim et al.²⁵⁰ showed that **SPy-EDTA** (Figure 7, 60) complexed with different paramagnetic metal ions enables structural studies of proteins by paramagnetic NMR. Because of the different enantiomeric forms of EDTA–metal complexes, the tag is suited for PRE measurements but is less so for PCS measurements, as demonstrated in a structure refinement of the protein domain ArgN using the **SPy-EDTA-Cu(II)** complex.²⁴⁹ Nonetheless, **SPy-EDTA** loaded with different lanthanoid ions proved successful in aligning a helical transmembrane protein for RDC measurements.⁶⁷⁵ As already discussed in section 5.2, **SPy-EDTA** loaded with Mn(II)⁶⁴⁴ and Cu(II)⁶⁴³ was also successfully applied to probe the structure of GB1 by solid-state NMR. Using solid-state NMR, Wang et al. identified glucuronoarabinoxylan, which is the major matrix polysaccharide in maize cell walls, as the binding target of β -expansins, by tagging the protein with **Mn(II)-SPy-EDTA** and measuring ^1H and ^{13}C PREs in ^{13}C enriched maize cell walls.⁶⁷⁶

4MMPyMTA (Figure 7, 73), **4PS-PyMTA** (Figure 7, 76), and **4PS-BrPyMTA** (Figure 7, 77) are three other frequently used probes employed in protein NMR research. **4PS-PyMTA** and **4PS-BrPyMTA** were used for elucidating the 3D structure of a protein in cells⁶¹⁰ and in pure protein solution (section 5.1.2).²⁶⁶ Su and co-workers labeled SrtA with **4PS-BrPyMTA** to assign NMR signals of an unstable intermediate by using PCSs and residue-selective isotope labeling.⁶⁷⁷ **4MMPyMTA** was used to detect the excited state of a protein–peptide complex by the new PCS-CEST method (section 5.1.5),²⁶³ and this approach was recently extended to ^{19}F NMR, referred to as ^{19}F PCS-CEST, to gain PCS restraints for protein–ligand interactions in the intermediate exchange regime.⁶⁷⁸ In a similar vein, ^1H PCSs of a protein–ligand complex presenting a minor species in solution were extracted from relaxation dispersion experiments (^1H PCS-RD experiments) to gain insight into the binding mode of a ligand to a bromodomain labeled with a **4MMPyMTA-Tm(III)** tag.⁶⁷⁹ The approach elegantly circumvents excessive line broadening in the intermediate exchange regime by detecting the PCSs of the minor species in the NMR signals of the ligand added in large excess.⁶⁷⁹

6.3. Applications of Cyclen-Based Probes

Section 5 gave some examples of applications of cyclen based probes in protein studies. Further examples are summarized below.

6.3.1. Double-Anchored Probes. The encounter complex between *Cc* and bovine adrenodoxin (Adx) was characterized by measuring PCSs and RDCs generated by the double-arm probe **Yb(III)-CLaNP-5** (Figure 9, 79) on *Cc*.⁷⁶ Loaded with different lanthanoid ions, **CLaNP-5** also delivered highly informative PCSs and PREs to model the structure of the little soluble 65

kDa complex between membrane associated Adx and adrenodoxin reductase (AdR).⁶⁸⁰

Saio et al. used **Yb(III)-CLaNP-5** to explore ligand-driven conformational changes of the multidomain protein MurD, where the PCSs identified a novel semiclosed conformation of the protein.⁶¹⁷ Venkata et al. used PREs generated by **Gd(III)-CLaNP-5** and **MTSL** to provide compelling evidence for two fundamentally different orientations of polyproline helices when bound to the Src SH3 domain, which are populated to different degrees.⁶⁸¹

The **CLaNP-5** tag has also proven useful to investigate conformational changes of proteins. In a maximum occurrence (MO) analysis of the different relative domain orientations of CaM, PCS, and PRE data generated by **Ln(III)-CLaNP-5** tags attached to either domain of CaM significantly improved the definition of the conformational space populated by the protein.⁶⁸² This tag was also used to study domain orientation in matrix metalloproteinase-1.⁶¹⁶ More recently, **Yb(III)-CLaNP-5** was used to investigate the conformational states of the single-domain GH11 glycosidase from *Bacillus circulans* (BCX), which proved to be in full agreement with crystal structures⁶⁸³ and only locally affected by site-directed mutagenesis.⁶⁸⁴

The double-anchored probe **CLaNP-7** (Figure 9, 80) proved similarly useful for the determination of the structure and encounter state of the complex between cytP450cam and putidaredoxin (Pdx), as discussed in section 5.1.3.^{629,631} Paramagnetic data recorded with the **Yb(III)-CLaNP-7** tag further demonstrated that the structure of the cytP450cam–Pdx complex predominantly populates a closed conformation in solution in contradiction to the open state reported by other crystal structures.⁶⁸⁵ **CLaNP-7** also proved useful to generate paramagnetic structure restraints of the heat-shock protein 90 (Hsp90) in the presence of proteins bound to the chaperone.^{686,687}

Except for a recent study on domain motions in diubiquitin,⁶¹⁴ the use of other double-arm probes, such as **T1/T2** (Figure 9, 82)⁶⁸⁸ and **CLaNP-13** (Figure 9, 85)²⁸¹ has only been explored for DEER distance measurements between two Gd(III) ions, with the aim to obtain narrow distance distributions by restricting the flexibility of the tags. In either case, the widths of the distance distributions obtained was not narrower than those obtained with some of the singly linked probes.

6.3.2. Applications of Single-Arm Cyclen Probes.

Among the single-arm cyclen based probes, **DOTA-M8** (Figure 9, 86) and **C1/C2** (Figure 9, 96) have been applied most frequently for protein structure investigations. The probe **C2** loaded with Dy(III) and Yb(III) attached at different sites was used to assign the methyl resonances of a four-helix bundle SNARE complex by measuring PCSs in ¹³C-HMQC spectra.⁶⁸⁹ PCSs induced in the C₂B domain of synaptotagmin-1 bound to the SNARE complex provided clear evidence for a highly dynamic binding mode between the two components.⁶⁹⁰

The dengue virus serotype 2 NS2B-NS3 protease (NS2B-NS3pro) is a recognized drug target but yields an overlapped and difficult to assign NMR spectrum. The interaction of NS2B with NS3pro was investigated by PCSs generated with probes **C1** and **C2** loaded with Tm(III) and Tb(III). The PCSs revealed that the enzyme predominantly assumes a closed conformation that is stabilized by the presence of positively charged active-site inhibitors.^{291,618} The same probes were used to investigate a NS2B-NS3pro construct without covalent

linkage, which confirmed the closed conformation in the absence of inhibitors.⁶¹⁹ A similar situation was documented for a linked construct of the related Zika virus NS2B-NS3 protease, where PCSs were induced in NS2B by **Tb(III)-C2** and **Tm(III)-C2** attached to NS3pro, revealing the closed conformation and stabilization by the presence of inhibitor.⁶⁹¹

PCSs generated by the **C1** and **C2** tags also provided the basis for the development of algorithms to determine the 3D structure of proteins, using the PCSs of backbone amides and backbone chemical shifts as the sole experimental restraints.^{94,604,606,612} Specifically, Pearce et al. demonstrated that two tagging sites can suffice to structurally define large segments of a protein, provided PCSs are available for multiple homo- and heteronuclear spins of the polypeptide backbone.⁹⁴ Using the combined information from PCSs and RDCs, the same study concluded that the molecular dynamics ensemble representation of ubiquitin (PDB ID 2K0X)⁶⁹² fitted the paramagnetic NMR data better than any single 3D structure available of the protein. Finally, the study showed that ¹H PCSs from tags at two sites in many cases suffices to determine the side-chain conformations of amino acids with methyl groups. In general, however, a complete 3D structure determination of a protein requires at least three, if not four, different probe tagging sites.

The **C1** tag loaded with Yb(III) and Tb(III) ions was also used to assess the binding mode of ¹⁵N/¹³C-labeled dGMP to the nucleotide binding R3H domain from human *Sμbp-2*, as well as the structural changes in the R3H domain in response to nucleotide binding.⁶⁹³ As an alternative to isotope labeling, a chemical tag, such as a *tert*-butyl group, also enables the selective detection of ¹H NMR signals of stably bound ligands and has been used to characterize the binding mode of a ligand to dengue virus NS2B-NS3pro by PCSs generated with **C2** loaded with Tm(III) and Tb(III) and detected in NOESY spectra.⁶³⁵

The **DOTA-M8** tag (Figure 9, 86) has been used in a particularly diverse range of applications, including the improvement of NMR spectral resolution of intrinsically disordered proteins (IDP),⁶⁹⁴ generation of RDCs to elucidate the structure of a transmembrane helix⁶⁹⁵ and an α -helix from myosin-VI that is remarkably stable in solution as a single helix,⁶⁹⁶ generation of RDCs in the ribosomal stalk protein bL12 to probe the orientational independence of N- and C-terminal domains,⁶⁹⁷ determining the fluorine coordinates of ligands bound to hCA II by PCSs measured in ¹⁹F NMR spectra recorded with four different tag attachment sites (including PCSs generated with **M7PyThiazole-DOTA**),⁶⁹⁸ and generation of PCSs in RNA.⁶⁹⁹ Specifically, Göbl et al. demonstrated that the relatively weak paramagnetism of Yb(III) in **DOTA-M8** was sufficient to improve spectral resolution in two different IDPs and enable additional resonance assignments,⁶⁹⁴ whereas Chiliveri et al. used **Tm(III)-DOTA-M8** to generate sizable molecular alignment of the HIV-1 gp41 transmembrane helix.⁶⁹⁵ Similarly, Wang et al. used **Tm(III)-DOTA-M8** to assess the domain alignment of ribosome-bound bL12 monomers and dimers by RDC measurements,⁶⁹⁷ Zimmermann et al. used the **Tm(III)-DOTA-M8** tag to generate PCSs in ¹⁹F NMR spectra of fluorine containing ligands bound to hCA II,⁶⁹⁸ and Strickland et al. showed that the RNA-binding protein U1A tagged with **Tm(III)-DOTA-M8** is capable of generating sizable PCSs in an RNA hairpin equipped with a U1A binding loop, which would otherwise be difficult to label with a paramagnetic tag.⁶⁹⁹

M7PyThiazole-DOTA (Figure 9, 92) is a newer probe than **DOTA-M8**, which offers a more rigid tether between lanthanide

ion and protein backbone. Loaded with Tm(III) and attached to each monomer of the homotrimeric protein Skp, a single set of PCSs was generated by M7PyThiazole-DOTA, which could be fitted by $\Delta\chi$ tensors positioned at three sites related by C_3 symmetry.⁷⁰⁰ Using DOTA-M8 and M7PyThiazole-DOTA as paramagnetic tags, Cucuzza et al. developed a new iterative procedure for structure refinement of repeat proteins based on backbone amide PCSs. The authors successfully determined the fold of the protein YM₄A in solution.⁷⁰¹ The related probes M7Py-DOTA (section 5.1.2)²⁸⁶ and M8-CAM-I,²⁸⁵ which likewise generate stable thioether linkages with cysteine, have been deployed for in-cell measurements of PCSs to obtain structural restraints with Tm(III) and Dy(III) ions, respectively, in GB1 and ubiquitin.

Lanthanoid tags can be deployed in DEER experiments when loaded with Gd(III), while generating PCSs in NMR experiments when loaded with other paramagnetic lanthanoid ions. Both approaches were combined by Abdelkader et al. in a structural analysis of the *E. coli* aspartate/glutamate binding protein with the C3 tag ligated to AzF.⁷⁰² Multiple sites tagged pairwise with Gd(III)-C3 delivered accurate positions of the metal ion relative to the crystal structure of the protein, which subsequently allowed fitting $\Delta\chi$ tensors using PCSs measured with Tm(III)-C3 and Tb(III)-C3 in 5-parameter fits that relied on the metal coordinates derived from the DEER measurements. The approach is of interest for proteins of limited solubility, where resonance assignment by 3D NMR experiments is difficult but 2D NMR spectra are accessible of samples prepared with selective isotope labeling, which can be assigned by high-throughput site-directed mutagenesis.

Gd(III) complexes of the above-mentioned probes and other single arm probes have been explored for distance measurements by DEER experiments in multiple EPR studies.^{298,703–709} Specifically the probes DO3MA-3BrPy (Figure 9, 107) and C3 (Figure 9, 98) were deployed for distance measurements in proteins embedded in cells.^{708,709}

6.4. Applications of Small Chemical Probes

Compared with the aforementioned probes, small chemical based probes have been less frequently used in paramagnetic NMR of proteins, which can be attributed to the inconvenience of having to titrate with paramagnetic metal ions after the tagging reaction and the risk of populating alternative binding sites with stray metal ions, which do not bind at the intended site either due to weak binding affinities to the probe or an excess of metal ion. Among the small chemical Gd(III) probes, 4MMDPA (Figure 11, 116) and the IDA-SH (Figure 11, 126) tag have been used for DEER distance measurements in proteins.^{688,710}

6.5. Applications of Cosolute Paramagnetic Probes

Cosolute paramagnetic probes are popular tools for detecting solvent accessible protein surfaces. Petros et al. explored the use of TEMPOL and [Gd(III)(DTPA)]²⁻ for the identification of solvent exposure of ubiquitin and hen egg white lysozyme (HEWL) by double-quantum-filtered proton correlation spectra (DQF-COSY), concluding that the two probes preferentially interacted with hydrophobic amino acid side chains and carboxylate groups, respectively.³⁴⁵ Fesik et al. subsequently showed that longitudinal ¹H PREs generated by TEMPOL identified the solvent-exposed residues of ¹³C-labeled cyclosporin bound to cyclophilin.³⁴⁶ The same group confirmed this approach using a ¹³C-labeled FK-506 analog bound to FKBP.⁷¹¹ Molinari et al. concluded that TEMPOL makes no specific

interactions with bovine pancreatic trypsin inhibitor (BPTI), allowing interpretation of signal attenuations in homo- and heteronuclear NMR spectra in terms of solvent exposure.⁷¹² This was followed by a string of paramagnetic perturbation studies of a number of proteins, including Tendamistat,⁷¹³ MNEI,³⁴⁶ HEWL,^{347,355} and Sso7d (from *Sulfolobus solfataricus*),⁷¹⁴ using TEMPOL, as well as soluble Gd(III) complexes as the paramagnetic agents. Bernini et al. investigated protein aggregation by monitoring the attenuation of ¹H NMR signals of BPTI in the presence of TEMPOL and Gd(III)DTPA-BMA as a function of protein concentration.⁷¹⁵ TEMPOL-induced paramagnetic perturbation was shown to be broadly compatible with results obtained on the solvent exposure of aromatic amino acid side chains by photochemically induced dynamic nuclear polarization (photo-CIDNP) experiments.³⁴⁷ The introduction of Gd(III)DTPA-BMA provided an alternative uncharged cosolute probe for paramagnetic perturbation studies, which is less hydrophobic than TEMPOL and was shown to deliver better correlations between calculated and experimental solvent PREs³⁵⁰ but in the case of α -bungarotoxin was reported to still show uneven attenuations of [¹³C,¹H]-HSQC cross-peaks of C ^{α} H groups across the protein,⁷¹⁶ which were reproduced by the much larger probe Gd₂L7.⁷¹⁷ To identify the interaction surface of a protein–protein complex by comparing cross-peak attenuations of the complex versus those of the individual proteins, a charged probe like [Gd(III)(EDTA)]¹⁻ was shown to be sufficient.³⁵⁷ Almeida et al. further demonstrated that also the highly charged probes [Gd(III)(DOTP)]⁵⁻ and [Gd(III)-DOTAM]³⁺ allow probing for protein–protein interactions.⁷¹⁸ It has been noted, however, that Gd(III) complexes with free coordination sites, such as in Gd(III)(DO3A), can favor transient associations with carboxylate groups on the protein surface.³⁶⁶

Different algorithms have been developed to exploit PREs from paramagnetic cosolutes for determining the 3D structures of proteins³⁵² and protein complexes,⁷¹⁹ as well as gaining information on protein dynamics.³⁵³ Madl et al. selected Gd(III)DTPA-BMA to aid protein structure determination, using a “second-sphere interaction model” to translate PREs into distance restraints between the NMR-active nuclei and the protein surface.³⁵² This became the basis of a strategy to integrate solvent PRE data generated by Gd(III)DTPA-BMA into the program HADDOCK⁷²⁰ to elucidate the structure of a 150 kDa ternary protein complex. Gd(III)DTPA-BMA also provided useful PRE restraints for the structure determination of the homodimeric protein Sam68,⁷²¹ and the probe clearly demonstrated how farnesylation alters the solvent accessibility of the C-terminal polypeptide segment of peroxisomal biogenesis factor 19 (PEX19).⁷²² By measuring transverse PREs of amide protons with Gd(III)DTPA-BMA as a function of protein concentration and taking into account local molecular crowding as an impediment to translational diffusion, Jens Led and co-workers identified a method to distinguish specific from nonspecific contacts in weakly self-associating human growth hormone.⁷²³ To eliminate exchange effects arising from rapid relaxation of Gd(III)-coordinated hydration water, the group of Chun Tang developed Gd(III)TTHA-TMA as a cosolute probe without free coordination sites on the Gd(III) ion, which allowed determining the populations of open and closed conformations from solvent PREs by comparison with theoretical expectations based on crystal structures and molecular dynamics simulations.³⁵³

As discussed in section 3.3.5, the $[\text{Ln}(\text{III})(\text{DPA})_3]^{3-}$ probe can generate measurable PCSs and its Gd(III) complex can produce PREs. Ma et al. investigated the binding of $[\text{Ln}(\text{III})(\text{DPA})_3]^{3-}$ probes (Ln = Eu, Gd, Tb, Tm, or Yb) to multiple sites in HEWL, which had been identified in a crystal structure of the protein with $[\text{Eu}(\text{III})(\text{DPA})_3]^{3-}$.⁷²⁴ While the paramagnetic effects observed could not be fully accounted for by the crystal structure, an MD simulation reproduced the highest affinity binding site and correlated small PCS magnitudes with increased local protein dynamics. More recently, Swarbrick et al. used PCSs generated by $[\text{Tm}(\text{III})(\text{DPA})_3]^{3-}$ to improve the spectral resolution of NOESY spectra of lipopeptide polymyxin B (PMXB) and the peptide MSI-594 bound to micelles.⁷²⁵ The PCSs originated from transient association of the probe with the peptides, but were not used as structural restraints. Instead, the concentration of the probe was tuned to improve the spectral resolution while limiting PREs, and the structures were determined from trNOEs.

Finally, it has been pointed out that, at a fundamental NMR-technical level, paramagnetic solvent probes can be used to simplify NMR spectral interpretation. For example, Schwarzingger and co-workers showed that difference spectra calculated for HSQC or NOESY experiments recorded in the presence and absence of Gd(III)DTPA-BMA can be used to focus on signals with little accessibility to the probe.⁷²⁶ Similarly, Sattler and Fesik proposed the use of high concentrations of $[\text{Dy}(\text{III})(\text{DTPA})]^{2-}$ to obtain substantial chemical shift changes in NMR spectra.³⁵⁸

7. CONCLUSIONS AND PROSPECTS

The invention and further development of paramagnetic and, in particular, site-specific probes has opened a wide range of protein structure investigation methods both by NMR spectroscopy and EPR spectroscopy. Although nitroxide spin labels have long been used with great success for PRE measurements and in EPR studies, many more structure restraints beyond distances become accessible with paramagnetic metal ions associated with anisotropic molecular magnetic susceptibility tensors. Lanthanoid ions stand out for their intrinsically large $\Delta\chi$ tensors and synthetic chemists have worked hard to optimize lanthanoid binding probes. A number of broad conclusions can be drawn from comparing the performance of these probes. (i) Site-specific attachment is difficult to attain without one or more covalent bonds. (ii) Rigid attachment of a lanthanoid ion is important for obtaining useful $\Delta\chi$ tensors and more easily achieved by double-arm attachment. In the case of proteins, immobilization can be achieved by additional coordination by carboxylate groups of amino acid side chains. (iii) Probes synthesized with the lanthanoid ion in place are more user-friendly and less prone to artifacts than probes that need to be titrated with metal ions following ligation with the target molecule. Therefore, cyclen based probes continue to be attractive despite the multistep protocols of their synthesis.

Significant challenges remain. Most covalently binding probes target cysteine residues and thus are incompatible with proteins containing natural cysteine residues that are accessible and functionally important. To avoid the dependence on cysteine residues, schemes for site-specific incorporation of noncanonical amino acid capable of binding lanthanoid probes have been developed. A generally satisfying solution has not yet been found, however, because either the resulting linker to the target protein is quite long (as in the case of, for example, probes attached by click chemistry), the unnatural amino acid disrupts

the protein structure (as in the case of, for example, phosphoserine), or the amino acid causes protein precipitation upon metal binding (as in the case of the bipyridyl and 8-hydroxy-quinoline amino acids). An emerging solution may be offered by the site-specific incorporation of photocaged selenocysteine, which, following decaging by UV irradiation, delivers a site that is much more reactive toward alkylating probes than cysteine thiol groups.^{727,728} The potential of harnessing this strategy for attaching double-armed probes is clear but has not yet been explored. Also the incorporation of an unnatural amino acid containing a stable nitroxide radical has not yet been achieved in a way that would dethrone the MTSL tag as the most popular spin label, partly because nitroxides are sensitive to reduction in live cells and partly because no aminoacyl-tRNA synthetase is available that is specific for a nitroxide amino acid featuring a short and rigid linker between nitroxide group and protein backbone.

Moving from lanthanoid ions to transition metal ions reduces the size of complexation cages and enables good binding affinities with nitrogen ligands but abandons the large and varied range of paramagnetic effects afforded by different lanthanoid ions. Attachment modes that leave vacant coordination sites on the metal ion risk leading to metal mediated protein dimerization. While this has not been observed routinely, it is a distinct possibility for many of the very small probes and also when a transition metal ion binding site is engineered by a pair of histidine residues.

With the advent of completely rigid probes, exciting prospects are opened for high-accuracy protein structure analysis, including determining the 3D structures of minor, little populated protein states by PCSs. With chemically stable ligation chemistry, such studies will foreseeably be possible also in living cells. Therefore, finding new rigid chelators remains an important goal for the future.

Finally, paramagnetic metal probes are becoming increasingly popular in EPR spectroscopy for measuring metal–metal distances in the nanometer range, where the metal ions can be Gd(III), Mn(II), or Cu(II) ions.^{121,162,163,729} Both NMR and EPR are based on spin interactions, and it is only logical that paramagnetic systems bring these two fields closer together. The combination of these spectroscopies provides a more complete picture of the world of biological macromolecules.

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Notes

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Biographies

Qing Miao completed her Ph.D. under the supervision of Prof. Marcellus Ubbink and Dr. Mark Overhand in 2019 at Leiden University. During her Ph.D. studies, she focused on paramagnetic NMR probe design, synthesis, and applications. Currently, she works as a junior research fellow at Shaanxi University of Science & Technology and is interested in natural abundance isotope analysis research.

Christoph Nitsche studied chemistry and business administration and received his Ph.D. in medicinal chemistry from Heidelberg University in 2014. He worked as a Feodor Lynen Fellow at the Australian National University and as a Rising Star Fellow at the Free University of Berlin. In 2020, he was appointed at the Australian National University, where he is currently Senior Lecturer and ARC DECRA Fellow in the Research School of Chemistry. His research program in medicinal chemistry and chemical biology focuses on drug discovery against infectious diseases, biocompatible chemistry, and peptide and protein modification.

Henry Orton completed his Ph.D. in 2020 at the Australian National University with a focus on experiment and software development for paramagnetic relaxation in biomolecular NMR spectroscopy. He is currently a postdoctoral fellow with Professor Gottfried Otting with interests in ultrafast magic angle spinning of biological solids.

Mark Overhand obtained a M. Sc. (Dec. 1989) from Leiden University (oligosaccharide synthesis, supervisors G.J. Boons, G. van der Marel, and J.H. van Boom) and a Ph.D. (Jan. 1995) from the University of Virginia, US (natural product synthesis, advisor S.M. Hecht). He did a postdoc. with a Swiss stipend at the ETH in Switzerland in the group of D.S. Seebach (design and synthesis of novel beta-hexapeptide helix) and returned to Leiden where he remains (1996–present): postdoc, assistant prof. (UD), associate prof. (UHD). Stanford: sabbatical (summer 2004). Awards: Golden spatula 1986 (B. sc. research), best teacher faculty of science 2006, Leiden University.

Gottfried Otting studied chemistry at the universities of Heidelberg and Freiburg (Germany) and received his Ph.D. in protein NMR spectroscopy from the ETH-Zürich (Switzerland) in 1987. He continued at the ETH-Zürich as a research assistant until 1992, when he was appointed professor of Molecular Biophysics at the Karolinska Institute in Stockholm (Sweden). In 2002, he was appointed professor at the Research School of Chemistry of the Australian National University in Canberra (Australia) on a Federation Fellowship by the Australian Research Council (ARC). He currently is an ARC Laureate Fellow at the same place. His research program focuses on method developments for protein NMR spectroscopy, including techniques for

the production of protein samples with labels for structure analysis and drug discovery.

Marcellus Ubbink obtained a Ph.D. in Chemistry from Leiden University in 1994, working on paramagnetic properties of metalloproteins. During his postdoc research at the University of Cambridge (UK), he developed a method to determine the structure of two proteins using paramagnetic NMR, based in intermolecular paramagnetic effects caused by a heme group. He returned to Leiden University in 1997 as an assistant professor and was appointed to associate professor (2004) and full professor (2010). Since 2019, he is also Programme Director of the bachelor Life Science and Technology. His current research interests comprise protein–protein interactions, enzymology, protein evolution, and paramagnetic protein NMR spectroscopy.

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ABBREVIATIONS

2AP	2-aminopyridine
AdR	adrenodoxin reductase
ArgN	<i>Escherichia coli</i> arginine repressor
BCX	single-domain GH11 glycosidase from <i>Bacillus circulans</i>
BPTI	bovine pancreatic trypsin inhibitor
BpyAla	bipyridyl alanine
BuLi	butyllithium
CAP	catabolite gene activator protein
Cc	yeast iso-1-cytochrome <i>c</i>
CcP	yeast cytochrome <i>c</i> peroxidase
CCR	cross-correlated relaxation
CDI	<i>N,N'</i> -carbonyldiimidazole
CEST	chemical exchange saturation transfer
CLaNP	caged lanthanoid NMR probe
COSY	correlation spectroscopy
CPMG	Carr–Purcell–Meiboom–Gill
CSA	chemical shift anisotropy
CTAB	cetyltrimethylammonium bromide
CuAAC	copper-catalyzed azide–alkyne cycloaddition
Cyclen	1,4,7,10-tetraazacyclododecane
Cys-Ph-TAHA	cysteiny-phenyl-triaminohexaacetate
cytP450cam	cytochrome P450cam
DCM	dichloromethane
DD	dipole–dipole
DEER	double-electron–electron resonance
DIPEA	<i>N,N</i> -diisopropylethylamine
DMF	<i>N,N</i> -dimethylformamide
DMSO	dimethyl sulfoxide
DNA	desoxyribonucleic acid
DNP	dynamic nuclear polarization
DOTA	1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid
dPA	2,2'-dipicolylamine
DPA	dipicolinic acid
DPAP	2,2-dimethoxy-2-phenylaceto-phenone
DQF-COSY	double-quantum-filtered COSY

- R_2^{Curie} paramagnetic enhancement of the transverse relaxation rate due to Curie mechanism
- Ω Larmor frequency of the nuclear spin
- σ_{N} ^1H spin shift tensor originated by the dipolar field of the ^{15}N spin
- σ_{\uparrow} positive ^1H spin shift tensor originated by the dipolar field of the ^{15}N spin
- σ_{\downarrow} negative ^1H spin shift tensor originated by the dipolar field of the ^{15}N spin

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